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(FILE 'HOME' ENTERED AT 15:22:31 ON 02 FEB 2004) SET COST OFF FILE 'HCAPLUS' ENTERED AT 15:22:50 ON 02 FEB 2004 E ALBUMIN/CT 753 S E3 L1132 S E11 L2 E E47+ALL 80101 S E2+NT L3 E E33+ALL 566 S E3, E2 L425218 S E2+NT L5 L6 157881 S ?ALBUMIN? L7 181833 S L1-L6 $\Gamma8$ 2969 S BDNF OR BD NF 2881 S BRAIN DERIVED NEUROTROPHIC FACTOR L9 2883 S (BD OR BRAIN DERIVED) () (NF OR NEUROTROPHIC FACTOR) L10 E NEUROTROPHIC FACTOR/CT 141 S E10 L11L12 2554 S E26 E E25+ALL 789 S E3-E5 AND BRAIN DERIVED L13 L14679 S E12,E13 L15 3242 S E2+NT (L) BRAIN DERIVED L16 64 S L7 AND L8-L15 L17 19234 S INTERFERONALPHA OR ALPHAINTERFERON OR INTERFERONBETA OR BETAI E INTERFERON/CT 302 S E3-E19 L18 18390 S E85-E101 E INTERFERONS/CT E E3+ALL 18391 S E7, E6 (L) (ALPHA OR BETA) L20. 546 S L7 AND L17-L20 L21 L22 2340 S TIMP()(I OR 1) FILE 'REGISTRY' ENTERED AT 15:29:36 ON 02 FEB 2004 1 S 140208-24-8 L23 FILE 'HCAPLUS' ENTERED AT 15:30:37 ON 02 FEB 2004 2026 S L23 L24 L25 859 S TISSUE INHIBITOR (1W) METALLOPROTEINASE 1 27 S METALLOPROTEINASE INHIBITOR 1 L26 L27 651 S TIMP1 L28 12 S FIBROBLAST COLLAGENASE INHIBITOR L29 91 S L7 AND L22, L24-L28 L30 678 S L16, L21, L29 L31 9815 S IFNALPHA OR IFNBETA OR ALPHAIFN OR BETAIFN OR IFN(A) (ALPHA OR L32 119 S L7 AND L31 L33 700 S L30, L32 L34 62 S L33 AND (FUSION OR FUSE OR FUSED OR FUSES OR FUSING) L35 167 S L33 AND RECOMBIN? L36 44 S L33 AND CHIMER? L37 202 S L34-L36 E ROSEN C/AU 27 S E3, E4 L38 E ROSEN CRAIG/AU 625 S E3-E5 L39 E HASELTINE W/AU L40 302 S E3, E4, E7-E10

10 S L33 AND L38-L40

E HUMAN GENOME SCI/PA, CS

L41

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975 S E5-E37
             13 S L33 AND L42
L43
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             13 S L41, L43
L45
             13 S L44 AND L37
              9 S L45 AND (SHELFLIFE OR SHELF LIFE)
L46
              4 S L45 NOT L46
L47
                SEL DN AN 1 4
              2 S L47 NOT E1-E6
L48
             11 S L46, L48
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                SEL RN
                DEL SEL
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L50
          11933 S E9
                E E9+ALL
           3795 S E3, E4
L51
L52
              5 S L51 AND L33
             29 S L50 AND L33
L53
L54
             34 S L49, L52, L53
             27 S L54 AND ALBUMIN
L55
              7 S L54 NOT L55
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L57
            159 S L37 AND ALBUMIN
L58
            132 S L57 NOT L43-L49, L52-L56
L59
              6 S L58 AND L16
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              7 S L58 AND L29
            121 S L58 NOT L59, L60
L61
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L62
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             18 S L62 AND E1-E54
L63
L64
             29 S L49, L63 AND L1-L22, L24-L63
L65
             29 S L64 AND ?ALBUMIN?
             29 S L64 AND (INF? OR INTERFERON OR TIMP? OR NEUROTROPHIC?)
L66
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والمتيترة

FILE 'HCAPLUS' ENTERED AT 16:00:16 ON 02 FEB 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE COVERS 1907 - 2 Feb 2004 VOL 140 ISS 6 FILE LAST UPDATED: 1 Feb 2004 (20040201/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L66 ANSWER 1 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN AN 2003:571103 HCAPLUS
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DN 139:122690

ED Entered STN: 25 Jul 2003

TI Albumin fusion proteins for prolonged shelf-life of therapeutic proteins

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ΙN
     Ballance, David James; Turner, Andrew John; Rosen, Craig A.; Haseltine,
     William A.
PA
     Human Genome Sciences, Inc., USA; Delta Biotechnology Limited; Principia
     Pharmaceutical Corporation
     PCT Int. Appl., 598 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
IC
     ICM C12N
CC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3
FAN.CNT 2
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     PATENT NO.
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                      A2
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                                           WO 2002-US40891 20021223
     WO 2003060071
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             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
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             MR, NE, SN, TD, TG
PRAI US 2001-341811P
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     US 2002-420246P
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                            20021023
     US 2002-423623P
                       Ρ
                            20021105
     The present invention encompasses albumin fusion proteins. Many
AΒ
     therapeutic proteins in their native state or when recombinantly produced .
     are typically labile mols. exhibiting short shelf-lives, particularly when
     formulated in aqueous solns.; fusions of the therapeutic protein with human
     serum albumin have a longer serum half-life and/or stabilized activity in
     solution (or in a pharmaceutical composition) in vitro and/or in vivo than the
     corresponding unfused therapeutic mols. Thus, albumin fusion proteins are
     provided comprising granulocyte colony-stimulating factor, interleukin 2,
     parathormone, erythropoietin, interferon \beta, interferon \alpha2,
     interferon A/D hybrid, a single-chain insulin analog, growth hormone, and
     (7-36) GLP-1. Nucleic acid mols. encoding the albumin fusion proteins of
     the invention are also encompassed by the invention, as are vectors containing
     these nucleic acids, host cells transformed with these nucleic acids
     vectors, and methods of making the albumin fusion proteins of the
     invention and using these nucleic acids, vectors, and/or host cells.
     Addnl. the present invention encompasses pharmaceutical compns. comprising
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albumin fusion proteins and methods of treating or preventing diseases,

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robinson - 09 / 833041 disorders or conditions related to diabetes mellitus using albumin fusion proteins of the invention. albumin fusion therapeutic protein shelflife Animal cell line (293, recombinant expression host; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) (CHO, recombinant expression host; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Animal cell line (NSO, recombinant expression host; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiviral, T1249 peptide inhibitor derived from HIV-1; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Antidiabetic agents Human Linking agents Molecular cloning (human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Fusion proteins (chimeric proteins) Interleukin 2 RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Signal peptides RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses) (human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Animal cell (mammalian, recombinant expression host; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Diabetes mellitus (non-insulin-dependent, treatment of; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Protein sequences (of human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Plasmid vectors (pC4; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Plasmid vectors (pEE12.1; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Plasmid vectors (pSAC35; human serum albumin fusion proteins for prolonged shelf-life of therapeutic proteins) Saccharomyces cerevisiae Yeast (recombinant expression host that is glycosylation and protease-deficient; human serum albumin fusion proteins for prolonged

Albumins, biological studies IT RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

shelf-life of therapeutic proteins)

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(serum; human serum albumin fusion proteins for prolonged shelf-life of
        therapeutic proteins)
     Interferons
TΥ
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\alpha 2; human serum albumin fusion proteins for prolonged shelf-life
        of therapeutic proteins)
ΙΤ
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\alpha; human serum albumin fusion proteins for prolonged shelf-life
        of therapeutic proteins)
IT
     Interferons
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\alpha AD; human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (β; human serum albumin fusion proteins for prolonged shelf-life
        of therapeutic proteins)
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     562119-82-8P
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                                    562119-85-1DP, Albumin (human),
     subfragments, fusion products
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (amino acid sequence; human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
                                 9004-10-8P, Insulin, biological studies
ΙT
     9002-64-6P, Parathormone
     11096-26-7P, Erythropoietin
                                   89750-14-1P, Glucagon-like peptide I
     143011-72-7P, Granulocyte colony-stimulating factor
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (human serum albumin fusion proteins for prolonged shelf-life of
        therapeutic proteins)
ΙT
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     RL: BUU (Biological use, unclassified); PRP (Properties); BIOL (Biological
     study); USES (Uses)
        (nucleotide sequence; human serum albumin fusion proteins for prolonged
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RL: PRP (Properties)
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        (unclaimed protein sequence; albumin fusion proteins for prolonged
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shelf-life of therapeutic proteins)

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     RL: PRP (Properties)
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DN
     139:122689.
     Entered STN: 25 Jul 2003
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     Albumin fusion proteins for prolonged shelf-
ΤI
     life of therapeutic proteins
ΙN
     Rosen, Craig A.; Haseltine, William A.
PΑ
     Human Genome Sciences, Inc., USA
SO
     PCT Int. Appl., 1086 pp.
     CODEN: PIXXD2
\mathsf{D}\mathbf{T}
     Patent
     English
LA
IC
     ICM C07K
CC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3
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                      KIND DATE
                                            APPLICATION NO.
                                                              DATE
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    US 2002-420246P
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    US 2002-423623P
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    The present invention encompasses albumin fusion
AΒ
    proteins. Many therapeutic proteins in their native state or when
    recombinantly produced are typically labile mols. exhibiting short
    shelf-lives, particularly when formulated in aqueous solns.;
    fusions of the therapeutic protein with human serum
    albumin have a longer serum half-life and/or stabilized activity
    in solution (or in a pharmaceutical composition) in vitro and/or in vivo than
the
    corresponding unfused therapeutic mols. Thus, albumin
    fusion proteins are provided comprising interferon .
    beta., interferon \alpha 2, insulin, bone
    morphogenetic protein 9, glucagon-like peptide-I(7-36), a hybrid
    interferon A/D, and extendin 4. Nucleic acid mols. encoding the
    albumin fusion proteins of the invention are also
    encompassed by the invention, as are vectors containing these nucleic acids,
    host cells transformed with these nucleic acids vectors, and methods of
    making the albumin fusion proteins of the invention
     and using these nucleic acids, vectors, and/or host cells. Addnl. the
    present invention encompasses pharmaceutical compns. comprising
     albumin fusion proteins and methods of treating or
    preventing diseases, disorders or conditions related to diabetes mellitus
    using albumin fusion proteins of the invention.
ST
    albumin fusion therapeutic protein shelflife
ΙT
    Animal cell line
        (293, recombinant expression host; human serum
        albumin fusion proteins for prolonged shelf
        -life of therapeutic proteins)
IT
    Animal cell line
        (CHO, recombinant expression host; human serum
        albumin fusion proteins for prolonged shelf
        -life of therapeutic proteins)
ΙT
    Animal cell line
        (NSO, recombinant expression host; human serum
        albumin fusion proteins for prolonged shelf
        -life of therapeutic proteins)
ΙT
    Metabolism, animal
        (disorder, treatment of; human serum albumin fusion
        proteins for prolonged shelf-life of therapeutic
        proteins)
ΤТ
    Antidiabetic agents
    Antiobesity agents
    Cardiovascular agents
    Human
    Linking agents
    Molecular cloning
        (human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
ΙT
     Fusion proteins (chimeric proteins)
    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
ΙT
     Signal peptides
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
     Diabetes mellitus
ΙT
        (insulin-dependent, treatment of; human serum albumin
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fusion proteins for prolonged shelf-life of

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therapeutic proteins)
ΙT
    Animal cell
        (mammalian, recombinant expression host; human serum
        albumin fusion proteins for prolonged shelf
        -life of therapeutic proteins)
ΙT
     Nerve, disease
        (neuropathy, treatment of; human serum albumin fusion
        proteins for prolonged shelf-life of therapeutic
        proteins)
ΙT
     Diabetes mellitus
        (non-insulin-dependent, treatment of; human serum albumin
        fusion proteins for prolonged shelf-life of
        therapeutic proteins)
     Protein sequences
ΙT
        (of human serum albumin fusion proteins for
        prolonged shelf-life of therapeutic proteins)
     Plasmid vectors
TT
        (pC4; human serum albumin fusion proteins for
        prolonged shelf-life of therapeutic proteins)
     Plasmid vectors
ΙT
        (pEE12.1; human serum albumin fusion proteins for
        prolonged shelf-life of therapeutic proteins)
ΙT
     Plasmid vectors
        (pSAC35; human serum albumin fusion proteins for
        prolonged shelf-life of therapeutic proteins) .
ΙT
     Saccharomyces cerevisiae
     Yeast
        (recombinant expression host that is glycosylation and
        protease-deficient; human serum albumin fusion
        proteins for prolonged shelf-life of therapeutic
        proteins)
TT
     Eye, disease
        (retinopathy, treatment of; human serum albumin
        fusion proteins for prolonged shelf-life of
        therapeutic proteins)
    Albumins, biological studies
ΙT
    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (serum; human serum albumin fusion proteins for
        prolonged shelf-life of therapeutic proteins)
     Cardiovascular system, disease
ΙT
     Endocrine system, disease
     Heart, disease
     Hyperglycemia
    Kidney, disease
     Nervous system, disease
     Obesity
        (treatment of; human serum albumin fusion proteins
        for prolonged shelf-life of therapeutic proteins)
TΤ
     Interferons
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\alpha 2; human serum albumin
        fusion proteins for prolonged shelf-life of
        therapeutic proteins)
IT
     Interferons
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\alpha ; human serum albumin fusion
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proteins for prolonged shelf-life of therapeutic

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proteins)
TT
    Interferons
    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (α AD; human serum albumin
        fusion proteins for prolonged shelf-life of
        therapeutic proteins)
IT
    Interferons
    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (\beta ; human serum albumin fusion
        proteins for prolonged shelf-life of therapeutic
       proteins)
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        (Unclaimed; albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
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    561347-54-4DP, Albumin (human), subfragments, fusion
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    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (amino acid sequence; human serum albumin fusion
        proteins for prolonged shelf-life of therapeutic
       proteins)
IT
     9004-10-8P, Insulin, biological studies
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    RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (human serum albumin fusion proteins for prolonged
        shelf-life of therapeutic proteins)
TΤ
     50-99-7, D-Glucose, biological studies
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (maintenance of basel level of; human serum albumin
        fusion proteins for prolonged shelf-life of
        therapeutic proteins)
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    study); USES (Uses)
        (nucleotide sequence; human serum albumin fusion
       .proteins for prolonged shelf-life of therapeutic
       proteins)
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561351-06-2

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        proteins for prolonged shelf-life of therapeutic
        proteins)
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                    561354-65-2
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     561354-79-8
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                                   561354-81-2
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                    561354-85-6
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                    561354-96-9
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     RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
        proteins for prolonged shelf-life of therapeutic
        proteins)
ΙT
     561350-49-0
                    561350-50-3
                                   561350-51-4
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561350-54-7
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                                            561352-65-6
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               561352-72-5
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                                            561352-76-9
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561352-79-2
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               561353-73-9
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                             561353-79-5
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                                                           561353-81-9
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                             561353-84-2
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561353-87-5
               561353-89-7
                             561353-90-0
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                                                           561353-92-2
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               561353-94-4
                             561353-95-5
                                            561353-96-6
                                                           561353-97-7
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561353-98-8
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561354-08-3
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                                                           561354-95-8
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RL: PRP (Properties)
   (unclaimed protein sequence; albumin fusion
   proteins for prolonged shelf-life of therapeutic
   proteins)
33017-11-7, Proinsulin C-peptide (human)
                                             40958-31-4, Somatostatin (sheep
          82177-09-1
                         85482-68-4
                                       85734-71-0
                                                    122024-47-9
reduced)
                             131748-18-0
                                            131748-19-1
                                                           157654-59-6
125677-89-6
               130912-02-6
166980-40-1
               170098-75-6
                             192503-43-8
                                            247166-37-6
                                                           367273-47-0
               477953-25-6
                             477953-26-7
                                            477953-27-8
                                                           477953-28-9
367273-48-1
               477.953-30-3
                             477953-31-4
                                            477953-32-5
                                                           477953-33-6
477953-29-0
               477953-35-8
                             478188-11-3
                                            478188-13-5
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                             561304-83-4
                                            561304-84-5
                                                           561304-85-6
               561304-87-8
                             561304-88-9
                                            561304-92-5
                                                           561304-95-8
561304-86-7
RL: PRP (Properties)
   (unclaimed sequence; albumin fusion proteins for
   prolonged shelf-life of therapeutic proteins)
ANSWER 3 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
2003:300832 HCAPLUS
138:326508
Entered STN:
              18 Apr 2003
Albumin fusion proteins with therapeutic proteins for
improved shelf-life
Rosen, Craig A.; Haseltine, William A.
Human Genome Sciences, Inc., USA
PCT Int. Appl., 457 pp.
CODEN: PIXXD2
Patent
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CC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3, 15
FAN.CNT 1
                                                            DATE
                     KIND DATE
                                           APPLICATION NO.
     PATENT NO.
                           -----
                                           _____
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                      ____
                                           WO 2002-US31794
                                                           20021004
PΙ
     WO 2003030821
                      A2
                            20030417
                      A3
                            20031211
     WO 2003030821
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD,
             RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
             NE, SN, TD, TG
                            20011005
PRAI US 2001-327281P
                      Ρ
     The present invention encompasses fusion proteins of
     albumin with various therapeutic proteins. Therapeutic proteins
     may be stabilized to extend the shelf-life, and/or to
     retain the therapeutic protein's activity for extended periods of time in
     solution, in vitro and/or in vivo, by genetically or chemical fusing
     or conjugating the therapeutic protein to albumin or a fragment
    or variant of albumin. Use of albumin fusion
     proteins may also reduce the need to formulate the protein solns. With
     large excesses of carrier proteins to prevent loss of therapeutic proteins
     due to factors such as binding to the container. Nucleic acid mols.
     encoding the albumin fusion proteins of the invention
     are also encompassed by the invention, as are vectors containing these nucleic
     acids, host cells transformed with these nucleic acids vectors, and
     methods of making the albumin fusion proteins of the
     invention and using these nucleic acids, vectors, and/or host cells.
     Thus, plasmid vectors are constructed in which DNA encoding the desired
     therapeutic protein may be inserted for expression of the albumin
     fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA).
     Yeast-derived signal sequences from Saccharomyces cerevisiae invertase
     SUC2 gene, or the stanniocalcin or native human serum albumin
     signal peptides, are used for secretion in yeast or mammalian systems,
     resp. Thus, the fusion product of human growth hormone with
     residues 1-387 of human serum albumin retains essentially intact
     biol. activity after 5 wk of incubation in tissue culture media at
     37°, whereas recombinant human growth hormone used as
     control lost its biol. activity in the first week. Although the potency
     of the albumin fusion proteins is slightly lower than
     the unfused counterparts in rapid bioassays, their biol. stability results
     in much higher biol. activity in the longer term in vitro assay or in vivo
             Addnl., the present invention encompasses pharmaceutical compns.
     Comprising albumin fusion proteins and methods of
     treating, preventing, or ameliorating diseases, disorders or conditions
     using albumin fusion proteins of the invention.
ST
     albumin fusion therapeutic protein shelflife
IT
     Drug delivery systems
     Gene therapy
     Human
     Molecular cloning
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
ΙT
     Fusion proteins (chimeric proteins)
       Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
ΙT
     Signal peptides
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
     Peptides, biological studies
IT
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (linkers; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT.
     Animal cell
        (mammalian, recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Plasmid vectors
IT
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Plasmid vectors
        (pScCHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
TT
     Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Linking agents
        (peptide; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Saccharomyces cerevisiae
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
TT
     Albumins, biological studies
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (serum; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Genetic element
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (signal sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
TT
     Antibodies
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (single chain; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IΤ
     Proteins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (therapeutic; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(\alpha ; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     9002-72-6DP, Growth hormone, fusion proteins with
ΙT
     albumin
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
     511566-72-6DP, Albumin (human blood serum), full-length or
ΙT
     subfragment fusion proteins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (amino acid sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     511566-73-7
     RL: BUU (Biological use, unclassified); PRP (Properties); BIOL (Biological
     study); USES (Uses)
        (nucleotide sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     511603-12-6
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                                 ·511603-14-8
                                                511603-15-9
                                                              511603-16-0
     511603-17-1
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                                                              511603-21-7
     511603-22-8
                   511603-23-9
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                                                511603-30-8
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     511603-67-1
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     122024-47-9
                                 367273-46-9
                                                367273-47-0
                                                              367273-48-1
IT
                   131748-18-0
     RL: PRP (Properties)
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     ANSWER 4 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
ΑN
     2003:125793 HCAPLUS
DN
     138:297265
ED
     Entered STN: 19 Feb 2003
TI
     An IFN-\beta -Albumin Fusion
     Protein That Displays Improved Pharmacokinetic and Pharmacodynamic
     Properties in Nonhuman Primates
ΑU
     Sung, Cynthia; Nardelli, Bernardetta; LaFleur, David W.; Blatter, Erich;
     Corcoran, Marta; Olsen, Henrik S.; Birse, Charles E.; Pickeral, Oxana K.;
     Zhang, Junli; Shah, Devanshi; Moody, Gordon; Gentz, Solange; Beebe, Lisa;
     Moore, Paul A.
CS
     Human Genome Sciences, Inc., Rockville, MD, 20850, USA
     Journal of Interferon and Cytokine Research (2003), 23(1), 25-36
     CODEN: JICRFJ; ISSN: 1079-9907
PB
     Mary Ann Liebert, Inc.
DT
     Journal
LA
     English
CC
     1-7 (Pharmacology)
     Section cross-reference(s): 15
     The long half-life and stability of human serum albumin (HSA)
AΒ
     make it an attractive candidate for fusion to short-lived
     therapeutic proteins. Albuferon beta (Human Genome Sciences [HGS], Inc.,
     Rockville, MD) is a novel recombinant protein derived from a
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 β ; IFN- β -albumin

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gene fusion of interferon-β
                             (
IFN-β ) and HSA. In vitro, Albuferon beta displays
antiviral and antiproliferative activities and triggers the IFN-stimulated
response element (ISRE) signal transduction pathway. Array anal. of 5694
independent genes in Daudi-treated cells revealed that Albuferon beta and
IFN-\beta induce the expression of an identical set of
.30 genes, including 9 previously not identified.
                                                  In rhesus monkeys
administered a dose of 50 µg/kg i.v. or s.c. or 300 µg/kg s.c.,
Albuferon beta demonstrated favorable pharmacokinetic properties. S.c.
bioavailability was 87%, plasma clearance at 4.7-5.7 mL/h/kg was approx.
140-fold lower than that of IFN-\beta , and the
terminal half-life was 36-40 h compared with 8 h for IFN-.
        Importantly, Albuferon beta induced sustained increases in
serum neopterin levels and 2',5'-oligoadenylate synthetase (2',5'-OAS)
mRNA expression. At a molar dose equivalent to one-half the dose of
IFN-β , Albuferon beta elicited comparable neopterin
responses and significantly higher 2',5'-OAS mRNA levels in rhesus
monkeys. The enhanced in vivo pharmacol. properties of IFN-.
beta. when fused to serum albumin suggest a
clin. opportunity for improved IFN-\beta therapy.
interferon beta albumin fusion
protein albuferon beta pharmacokinetic pharmacodynamic
Fusion proteins (chimeric proteins)
RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PKT
(Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
    (IFN-\beta -HSA; IFN-\beta -
   albumin fusion protein with retained biol. activities
   and improved pharmacokinetic and pharmacodynamic properties of
   IFN-\beta in primates)
Antiviral agents
Human
Macaca mulatta
Pharmacodynamics
Pharmacokinetics
Signal transduction, biological
    (IFN-\beta -albumin fusion
   protein with retained biol. activities and improved pharmacokinetic and
   pharmacodynamic properties of IFN-\beta in
   primates)
Genetic element
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (ISRE (interferon-stimulated response element); IFN
   -β -albumin fusion protein with
   retained biol. activities and improved pharmacokinetic and
   pharmacodynamic properties of IFN-\beta in
   primates)
Transcriptional regulation
   (activation; IFN-\beta -albumin
   fusion protein with retained biol. activities and improved
   pharmacokinetic and pharmacodynamic properties of IFN-
   β in primates)
Cell proliferation
   (inhibition; IFN-\beta -albumin
   fusion protein with retained biol. activities and improved
   pharmacokinetic and pharmacodynamic properties of IFN-
      in primates)
Albumins, biological studies
RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PKT
(Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
    (serum, human, fusion protein with IFN-
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robinson - 09 / 833041 fusion protein with retained biol. activities and improved pharmacokinetic and pharmacodynamic properties of IFNin primates) Interferons RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (β , fusion protein with albumin; IFN- β -albumin fusion protein with retained biol. activities and improved pharmacokinetic and pharmacodynamic properties of $IFN-\beta$ in primates) 507485-69-0P, Albuferon beta RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (IFN- β -HSA; IFN- β albumin fusion protein with retained biol. activities and improved pharmacokinetic and pharmacodynamic properties of **IFN-** β in primates) 69106-44-1, 2',5'-Oligoadenylate synthetase 2009-64-5, Neopterin RL: BSU (Biological study, unclassified); BIOL (Biological study) (IFN- β -albumin fusion protein with retained biol. activities and improved pharmacokinetic and pharmacodynamic properties of IFN- β primates) THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT (1) Brumell, J; J Immunol 1999, V163, P3388 HCAPLUS (2) Chuang, V; Pharm Res 2002, V19, P569 (3) Durelli, L; Lancet 2002, V359, P1453 HCAPLUS (4) Eisen, M; Proc Natl Acad Sci USA 1998, V95, P14863 HCAPLUS (5) Fierlbeck, G; J Interferon Cytokine Res 1996, V16, P777 MEDLINE (6) Fine, H; Clin Cancer Res 1997, V3, P381 HCAPLUS (7) Fukutomo, T; J Hepatol 2001, V34, P100 (8) Glue, P; Clin Pharmacol Ther 2000, V68, P556 HCAPLUS (9) Grace, M; J Interferon Cytokine Res 2001, V21, P1103 HCAPLUS (10) Gutterman, J; Proc Natl Acad Sci USA 1994, V91, P1198 HCAPLUS (11) Imaizumi, T; J Leukocyte Biol 2002, V72, P486 HCAPLUS (12) Jacobs, L; N Engl J Med 2000, V343, P898 HCAPLUS (13) Karsan, A; Blood 1996, V87, P3089 HCAPLUS (14) Kho, C; J Biol Chem 1997, V272, P13426 HCAPLUS (15) Lafleur, D; J Biol Chem 2001, V276, P39765 HCAPLUS (16) Leaman, D; J Biol Chem 2002, V277, P28504 HCAPLUS (17) Lindsay, K; Hepatology 2001, V34, P395 HCAPLUS (18) Lukashok, S; J Virol 2000, V74, P4705 HCAPLUS (19) Maeyer, E; The Cytokine Handbook, 3rd ed 1998, P491 (20) Marques, J; Thromb Haemost 2001, V86, P902 HCAPLUS (21) Osborn, B; Eur J Pharmacol 2002, V456, P149 HCAPLUS (22) Osborn, B; J Pharmacol Exp Ther 2002, V303, P540 HCAPLUS (23) Paty, D; Neurology 1993, V43, P662 MEDLINE (24) Pellegrini, S; Mol Cell Biol 1989, V9, P4605 HCAPLUS (25) Pepinsky, R; J Pharmacol Exp Ther 2001, V297, P1059 HCAPLUS (26) Peters, T; All About Albumin 1996 (27) Pferrer, L; Cancer Res 1998, V58, P2489 (28) Prisms Study Group; Lancet 1998, V352, P1498 (29) Prisms Study Group and the University of British Columbia MS/MRI Analysis Group; Neurology 2001, V56, P1628 (30) Runkel, L; Pharm Res 1998, V15, P641 HCAPLUS (31) Salmon, P; J Interferon Cytokine Res 1996, V16, P759 HCAPLUS

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- ANSWER 5 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN L66
- 2002:834389 HCAPLUS ΑN
- DN 137:304506
- Entered STN: 03 Nov 2002 ΕD
- Pharmacokinetic and pharmacodynamic studies of a human serum ΤI albumin-interferon- α fusion protein in cynomolgus monkeys
- Osborn, Blaire L.; Olsen, Henrik S.; Nardelli, Bernardetta; Murray, James AU H.; Zhou, Joe X. H.; Garcia, Andrew; Moody, Gordon; Zaritskaya, Liubov S.; Sung, Cynthia
- CS Human Genome Sciences, Inc., Rockville, MD, USA
- Journal of Pharmacology and Experimental Therapeutics (2002), 303(2), SO CODEN: JPETAB; ISSN: 0022-3565
- American Society for Pharmacology and Experimental Therapeutics PΒ
- DTJournal
- LA English
- 1-7 (Pharmacology) CCSection cross-reference(s): 15
- AB Interferon- α (IFN- α)

is indicated for the treatment of certain viral infections including hepatitis B and C, and cancers such as melanoma. The short circulating half-life of unmodified IFN- α makes frequent dosing (daily or three times weekly) over an extended period (6-12 mo or more) necessary. To improve the pharmacokinetics of IFN- α and decrease dosing frequency, IFN

was fused to human serum albumin

producing a new protein, Albuferon. In vitro comparisons of Albuferon and IFN- α showed similar antiviral and

antiproliferative activities, although Albuferon was less potent on a molar basis than ${\tt IFN-}\alpha$. Pharmacokinetic and pharmacodynamic properties of the fusion protein were enhanced

in monkeys. After a single i.v. injection (30 μ g/kg) clearance was 0.9 mL/h/kg, and the terminal half-life was 68 h. After 30 $\mu g/kg$ s.c. injection, apparent clearance (clearance divided by bioavailability) was 1.4 mL/h/kg, the terminal half-life was 93 h, and bioavailability was 64%. The rate of clearance of Albuferon was approx. 140-fold slower, and the half-life 18-fold longer, than for IFN- α given

by the s.c. route in other monkey studies. Sera from Albuferon-treated monkeys demonstrated dose-related antiviral activity for ≥8 days based on an in vitro bioassay, whereas antiviral activity from IFN $-\alpha$ -treated animals was only slightly elevated relative to vehicle on day 0. Significant increases in 2',5'-oligoadenylate

synthetase mRNA relative to $IFN-\alpha$ - or

vehicle-treated animals were maintained for ≥10 days after s.c. The improved pharmacokinetics of Albuferon are accompanied by an improved pharmacodynamic response suggesting that Albuferon may offer the benefits of less frequent dosing and a potentially improved efficacy profile compared with IFN- α .

- Albuferon interferon antiviral antiproliferative pharmacokinetics pharmacodynamics
- ΙT Antiviral agents Cytotoxic agents Human Macaca irus

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Pharmacodynamics
     Pharmacokinetics
        (pharmacokinetic and pharmacodynamic studies of a human serum
        albumin-interferon-\alpha fusion
        protein in cynomolgus monkeys)
ΙT
     Albumins, biological studies
     RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (serum, fusion protein with interferon-
        \alpha ; pharmacokinetic and pharmacodynamic studies of a human
        serum albumin-interferon-\alpha
        fusion protein in cynomolgus monkeys)
IT
     Interferons
     RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\alpha , fusion protein with human serum
        albumin; pharmacokinetic and pharmacodynamic studies of a human
        serum albumin-interferon-α
        fusion protein in cynomolgus monkeys)
     69106-44-1, 2',5'-Oligoadenylate synthetase
ΙT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (pharmacokinetic and pharmacodynamic studies of a human serum
        albumin-interferon-α fusion
        protein in cynomolgus.monkeys)
                            472960-22-8, Albuferon
ΙT
     98530-12-2, Intron-A
     RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (pharmacokinetic and pharmacodynamic studies of a human serum
        albumin-interferon-\alpha fusion
        protein in cynomolgus monkeys)
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L66
     2001:781112 HCAPLUS
ΑN
     135:348852
DN
     Entered STN: 26 Oct 2001
ED
ΤI
     Albumin fusion proteins with therapeutic proteins for
     improved shelf-life
     Rosen, Craig A.; Haseltine, William A.
ΙN
PΑ
     Human Genome Sciences, Inc., USA
SO
     PCT Int. Appl., 394 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
     ICM C12N015-00
IC
CC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3, 15
FAN.CNT 7
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
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                                           WO 2001-US11991
                                                            20010412
     WO 2001079480
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                      C2
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            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
             VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           EP 2001-937179
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                            20030122
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     EP 1276856
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     US 2003125247
                      A1
                            20030703
                                           US 2001-833041
                                                            20010412
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     US 2003171267
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                            20030911
                                           US 2001-833117
     JP 2003530852
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                            20031023
                                           US 2001-832501
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     US 2003199043
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                                           US 2001-833118
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                            20031127
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                                           US 2001-833245
     US 2004010134
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                            20040115
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PRAI US 2000-229358P
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     US 2000-199384P
                       Ρ
                            20000425
     US 2000-256931P
                      Ρ
                            20001221
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     WO 2001-US11991
                         . 20010412
     The present invention encompasses fusion proteins of
AB
     albumin with various therapeutic proteins. Therapeutic proteins
     may be stabilized to extend the shelf-life, and/or to
     retain the therapeutic protein's activity for extended periods of time in
     solution, in vitro and/or in vivo, by genetically or chemical fusing
     or conjugating the therapeutic protein to albumin or a fragment
     or variant of albumin. Use of albumin fusion
     proteins may also reduce the need to formulate the protein solns. with
     large excesses of carrier proteins to prevent loss of therapeutic proteins
     due to factors such as binding to the container. Nucleic acid mols.
     encoding the albumin fusion proteins of the invention
     are also encompassed by the invention, as are vectors containing these nucleic
     acids, host cells transformed with these nucleic acids vectors, and
     methods of making the albumin fusion proteins of the
     invention and using these nucleic acids, vectors, and/or host cells.
     Thus, plasmid vectors are constructed in which DNA encoding the desired
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therapeutic protein may be inserted for expression of the albumin fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA). Yeast-derived signal sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the stanniocalcin or native human serum albumin signal peptides, are used for secretion in yeast or mammalian systems, resp. Thus, the fusion product of human growth hormone with residues 1-387 of human serum albumin retains essentially intact biol. activity after 5 wk of incubation in tissue culture media at 37°, whereas recombinant human growth hormone used as control lost its biol. activity in the first week. Although the potency of the albumin fusion proteins is slightly lower than the unfused counterparts in rapid bioassays, their biol. stability results in much higher biol. activity in the longer term in vitro assay or in vivo Addnl,, the present invention encompasses pharmaceutical compns. comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention. albumin fusion therapeutic protein shelflife Receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (4-1BB; albumin fusion proteins with therapeutic proteins for improved shelf-life) Cytokines RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (BAFF; albumin fusion proteins with therapeutic proteins for improved **shelf-life**) Cytokine receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (DR4 (death receptor 4); albumin fusion proteins with therapeutic proteins for improved shelf-life) Cytokine receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (DR5 (death receptor 5); albumin fusion proteins with therapeutic proteins for improved shelf-life) Cytokines RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (MPIF-1 (myeloid progenitor inhibitory factor 1); albumin fusion proteins with therapeutic proteins for improved shelf-life) Steroid receptors Thyroid hormone receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (TR (thyroid/steroid hormone receptor), 11; albumin fusion proteins with therapeutic proteins for improved shelf-life) Steroid receptors Thyroid hormone receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (TR (thyroid/steroid hormone receptor), 12; albumin fusion proteins with therapeutic proteins for improved shelf-life) Steroid receptors Thyroid hormone receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(TR (thyroid/steroid hormone receptor), 13; albumin

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fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
    Steroid receptors
    Thyroid hormone receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TR (thyroid/steroid hormone receptor), 14; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Steroid receptors
    Thyroid hormone receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TR (thyroid/steroid hormone receptor), 16; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
    Steroid receptors
    Thyroid hormone receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TR (thyroid/steroid hormone receptor), 8; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
    Steroid receptors
ΙT
    Thyroid hormone receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TR2 (thyroid/steroid hormone receptor 2); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
     Steroid receptors
    Thyroid hormone receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TR3 (thyroid/steroid hormone receptor 3); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TRAIL (tumor necrosis factor-related apoptosis-inducing ligand);
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
    Cytokine receptors
ΙT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TRAIL, 4; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Cytokine receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TRAIL, 6; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Cytokine receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TRAIL-R3; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Drug delivery systems
    Gene therapy
    Molecular cloning
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
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مه ونيتوز

المرابعة

٠٠ تانيتنونه

٠ تايترل

ور ونيستان

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Cell adhesion molecules
TΨ
     Cytokines
     Enzymes, biological studies
     Fas antigen
     Fas ligand
       Fusion proteins (chimeric proteins)
     Growth factors, animal
       Interferons
     Synthetic gene
     Tumor necrosis factor receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
     Proteins, specific or class
TΤ
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (apoptosis-regulating, AIM-2; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Cytokines
ΙT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (endokine; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙΤ
     Signal peptides
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (for improved secretion in yeast or mammalian cells; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (keratinocyte-derived; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
TT
     Animal cell
        (mammalian, recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT.
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Plasmid vectors
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Plasmid vectors
        (pScCHSa, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
ΙT
     Saccharomyces cerevisiae
     Yeast
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Albumins, biological studies
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RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic

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use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (serum; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
     Genetic element
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
      (Uses)
         (signal sequence, for improved secretion in yeast or mammalian cells;
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
IT
     Antibodies
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (single chain; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
TI
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (therapeutic; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (\alpha ; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
· TT
     Chemokine receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (β chemokine receptor CCR5; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Tumor necrosis factors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
             albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
     Tumor necrosis factors
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (\delta; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
     189460-40-0P, Connective tissue growth factor
TΤ
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (2 and 4; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
     9001-84-7P, Phospholipase A2
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (T-cell lymphoma lipoprotein-associated; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
                                        9002-68-0P, FSH
                                                           9002-72-6P, Growth
ΙT
     9002-67-9P, Luteinizing hormone
                9004-10-8P, Insulin, biological studies
                                                          11096-26-7P,
     hormone
                       67763-96-6P, Insulin-like growth factor 1
                                                                   83869-56-1P,
     Erythropoietin
              124861-55-8P, Proteinase inhibitor, TIMP-2
     GM-CSF
     127464-60-2P, Vascular endothelial growth factor 140208-24-8P,
     Proteinase inhibitor, TIMP-1
                                     143011-72-7P, G-CSF
     145809-21-8P, Proteinase inhibitor, TIMP-3
                                                   148348-15-6P,
                                   171758-70-6P, Keratinocyte growth factor 2
     Fibroblast growth factor 7
     186207-03-4P, Proteinase inhibitor, TIMP-4
                                                   205944-50-9P,
                                                                244019-42-9P,
     Osteoprotegerin
                        207621-35-0P, Osteoprotegerin ligand
     Vascular endothelial growth factor 2
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
    127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A
ΙT
    protein moiety reduced), full-length or subfragment fusion
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (nucleotide sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
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    RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
       proteins with therapeutic proteins for improved shelf-
        life)
                  221879-28-3
                                222614-92-8
                                               352583-76-7, Protein (human
ΙT
    173586-11-3
     clone 785CIP2B 67)
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                                      370649-85-7
     RL: PRP (Properties)
        (unclaimed protein sequence; albumin fusion
       proteins with therapeutic proteins for improved shelf-
        life)
                                244008-03-5, PN: WO9947540 SEQID: 3 unclaimed
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    WO9947540 SEQID: 9 unclaimed DNA 244008-14-8, PN: WO9947540 SEQID: 10
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     370649-86-8
    RL: PRP (Properties)
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
             THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Delta Biotechnology Limited; EP 0322094 A1 1989 HCAPLUS
(2) Delta Biotechnology Limited; WO 9523857 A1 1995 HCAPLUS
    ANSWER 7 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
ΑN
     2001:781079 HCAPLUS
   . 135:348851
DN
    Entered STN: 26 Oct 2001
ED
    Albumin fusion proteins with therapeutic proteins for
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improved shelf-life
ΙN
    Rosen, Craig A.; Haseltine, William A.
PA
     Human Genome Sciences, Inc, USA
SO
     PCT Int. Appl., 606 pp.
     CODEN: PIXXD2
DT
     Patent
LA
    English
IC
     ICM C12N
CC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3, 15
FAN.CNT 7
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO. DATE
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    WO 2001079444
                      A2
                            20011025
                                           WO 2001-US12013 20010412
PΙ
    WO 2001079444
                     AЗ
                            20020523
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             CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
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             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
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     AU 2001074809
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                                         AU 2001-74809
                                                           20010412
     EP 1278544
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                            20030129
                                          EP 2001-941457 20010412
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     US 2003125247
                      A1
                            20030703
                                           US 2001-833041
                                                            20010412
     US 2003171267
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                                                            20010412
     JP 2003530847
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    US 2003199043
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    US 2003219875
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     US 2004010134
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PRAI US 2000-229358P
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                            20000412
    US 2000-199384P
                      Ρ
                            20000425
     US 2000-256931P
                      Ρ
                            20001221
    WO 2001-US12013
                     W
                            20010412
    The present invention encompasses fusion proteins of
AB
    albumin with various therapeutic proteins. Therapeutic proteins
    may be stabilized to extend the shelf-life, and/or to
     retain the therapeutic protein's activity for extended periods of time in
     solution, in vitro and/or in vivo, by genetically or chemical fusing
    or conjugating the therapeutic protein to albumin or a fragment
    or variant of albumin. Use of albumin fusion
    proteins may also reduce the need to formulate the protein solns. with
     large excesses of carrier proteins to prevent loss of therapeutic proteins
     due to factors such as binding to the container. Nucleic acid mols.
     encoding the albumin fusion proteins of the invention
    are also encompassed by the invention, as are vectors containing these nucleic
     acids, host cells transformed with these nucleic acids vectors, and
    methods of making the albumin fusion proteins of the
     invention and using these nucleic acids, vectors, and/or host cells.
     Thus, plasmid vectors are constructed in which DNA encoding the desired
     therapeutic protein may be inserted for expression of the albumin
    fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA).
     Yeast-derived signal sequences from Saccharomyces cerevisiae invertase
     SUC2 gene, or the stanniocalcin or native human serum albumin
     signal peptides, are used for secretion in yeast or mammalian systems,
    resp. Thus, the fusion product of human growth hormone with
    residues 1-387 of human serum albumin retains essentially intact
    biol. activity after 5 wk of incubation in tissue culture media at
     37°, whereas recombinant human growth hormone used as
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control lost its biol. activity in the first week. Although the potency of the albumin fusion proteins is slightly lower than the unfused counterparts in rapid bioassays, their biol. stability results in much higher biol. activity in the longer term in vitro assay or in vivo assays. Addnl., the present invention encompasses pharmaceutical compns. comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention. albumin fusion therapeutic protein shelflife Chemokines RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (1-309; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (11; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (12; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (15; albumin fusion proteins with therapeutic proteins for improved **shelf-life**) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (17; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (18; albumin fusion proteins with therapeutic proteins for improved shelf-life) Interleukins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (19; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (1; albumin fusion proteins with therapeutic proteins for improved shelf-life) Interleukins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (21; albumin fusion proteins with therapeutic proteins for improved shelf-life) Bone morphogenetic proteins RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (2; albumin fusion proteins with therapeutic proteins for improved shelf-life) Chemokines RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(331D5; albumin fusion proteins with therapeutic

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proteins for improved shelf-life)
    Bone morphogenetic proteins
ΤТ
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (3; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (4-1BB; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    Bone morphogenetic proteins
IT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (4; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Bone morphogenetic proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (5; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (61164; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Bone morphogenetic proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (6; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Bone morphogenetic proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (7; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
TΤ
    Bone morphogenetic proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (9; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Platelet-derived growth factors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (AA; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ACRP-30; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ADEC (adenoid expressed chemokine); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Interleukins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (AGF; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
     Proteins, specific or class
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RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (APM-1; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (Act-2; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Platelet-derived growth factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (BB; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (BCMA; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Platelet-derived growth factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (Bv-sis; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, 2; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, 3; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, DGWCC; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, DVic-1; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, ELC; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, HCC-1; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, IBICK; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (C-C, ILINCK; albumin fusion proteins with
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therapeutic proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C-C, SLC (secondary lymphoid chemokine); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C-C, STCP-1; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C-X-C, 3; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C-X-C; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
ΙT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C10; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
ΙT
    Troponins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (C; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CCC3; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); .PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CCF18; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CCR2; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
IT
    CD antigens
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CD27; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
IT
    Glycoproteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CD40-L (antigen CD40 ligand); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CTAP-III (connective tissue activating protein III); albumin
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fusion proteins with therapeutic proteins for improved

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shelf-life)
IT
    Antigens
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CTLA-8; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Chemokine receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (CXCR3; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
TΤ
     Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Cerebus; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Chr19Kine; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Platelet-derived growth factors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (D; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Cytokine receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (DR3 (death receptor 3); albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
     Proteins, specific or class
ΙT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (EDAR; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Interleukins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (EDIRF I protein; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
    Chemokines
IT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (EEC (eosinophil expressed chemokine); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ENA-78 (epithelial neutrophil activating protein-78); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
    Hemopoietins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (FLT3 ligand; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (HCC-1; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
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ΙT
    Troponins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (I; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (L105-7; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (LVEC-1 (liver expressed chemokine 1); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (LVEC-2 (liver expressed chemokine 2); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Lyn-1; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (M110; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (M11A; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (MACK (mammary associated chemokine); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (MCP-3\alpha and MCP-3\beta;
                              albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (MCP-4; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
TT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (MCPP (monocyte chemotactic proprotein); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(MDC (macrophage-derived chemokine); albumin fusion
   proteins with therapeutic proteins for improved shelf-
Monokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (MIG (monokine induced by \gamma- interferon);
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (MIG-\beta; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Interleukins
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (MIRAP; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (MP52; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (NOGO-66; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (NOGO-A; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (NOGO-B; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (NOGO-C; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Antigens
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (OX-40; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (PF4; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (PGBC (pituitary expressed chemokine); albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Chemokine receptors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(RANTES; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (SISD; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (SLC (secondary lymphoid tissue chemokine); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Troponins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (T; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TAC1; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Cytokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TARC (thymus and activation regulated cytokine); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TMEC (T cell mixed lymphocyte reaction expressed chemokine);
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Tarc; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Tim-1; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Troy; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
TT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ZCHEMO-8; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ZSIG-35; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Drug delivery systems
     Gene therapy
     Molecular cloning
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(albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
IT
    CD30 (antigen)
    CD40 (antigen)
    Cell adhesion molecules
    Cytokines
    Enzymes, biological studies
    Eotaxin
    Erythropoietin receptors
    Fas ligand
       Fusion proteins (chimeric proteins)
    Granulocyte-macrophage colony-stimulating factor receptors
    Growth factors, animal
       Interferons
    Interleukin 1
    Interleukin 1 receptor antagonist
    Interleukin 11
    Interleukin 13
    Interleukin 14
    Interleukin 15
    Interleukin 17
    Interleukin 18
    Interleukin 1\alpha
    Interleukin 1B
    Interleukin 3
    Interleukin 4
    Interleukin 4 receptors
    Interleukin 5 receptors
    Interleukin 6
    Interleukin 6 receptors
    Interleukin 8
    Interleukin 8 receptors
    Interleukin 9
    Lymphotoxin
    Monocyte chemoattractant protein-1
    Neutrophil-activating peptide-2
    Platelet-derived growth factors
    RANTES (chemokine)
    Stem cell factor
    Synthetic gene
    Tumor necrosis factor receptors
    Tumor necrosis factors
    Vascular endothelial growth factor receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
ΙT
    Interleukin 10
     Interleukin 12
    Interleukin 2
    Interleukin 5
    Interleukin 7
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
IT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (b57; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
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(chemokine-like protein PF4-414; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Growth factors, animal
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (chondromodulins, -like protein; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Proteins, specific or class
ΙT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (collapsins, antibodies for; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (exodus; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Signal peptides
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (for improved secretion in yeast or mammalian cells; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Chemokines
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (fractalkines; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Agglutinins and Lectins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (galectin-4; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gene Patched-2; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Vascular endothelial growth factor receptors
TΥ
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gene flt 1; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Vascular endothelial growth factor receptors
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gene flt 4; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Proteins, specific or class
ΙT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gene patched; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (glycodelin-A; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(granulocyte chemotactic protein-2; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gro-\alpha; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gro-\beta; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (gro-γ; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (growth-related oncogene-\alpha; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (growth-related oncogene-\beta; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (growth-related oncogene-γ; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Cytokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interferon-inducible IP-10; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Interleukin receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interleukin 10 receptors; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
     Interleukin receptors
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interleukin 11; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Interleukin receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interleukin 12; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Interleukin receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interleukin 13; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Interleukin receptors
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RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin 15; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Interleukin receptors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin 17; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Interleukin receptors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin 9; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin C; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin-1 accessory; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (interleukin-2 receptor associated p43; albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Lymphokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (lymphotactins; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (macrophage inflammatory protein 3\alpha; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (macrophage inflammatory protein 3\beta; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (macrophage inflammatory protein 3\gamma; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Animal cell
   (mammalian, recombinant expression host; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Antitumor agents
   (melanoma; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Chemokines
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(monocyte chemoattractant protein 3; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Chemokine receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (monocyte chemoattractant protein-1; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (monocyte chemoattractant protein-2; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Chemokine receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (monocyte chemoattractant protein-4; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (neurotactin; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Growth factors, animal
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (osteogenic protein 2; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Tumor necrosis factor receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (p75; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
     Plasmid vectors
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Plasmid vectors
        (pScCHSa, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
IΤ
     Placental hormones
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (placenta-derived mitogenic factors; albumin fusion
        proteins with therapeutic proteins for improved shelf-
ΙT
     Saccharomyces cerevisiae ·
     Yeast
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
```

shelf-life)

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Albumins, biological studies
ΙT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (serum; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
TT
    Genetic element
    RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (signal sequence, for improved secretion in yeast or mammalian cells;
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
ΙT
    Antibodies
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (single chain; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Chemokines
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (stem cell inhibitory factor; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Growth factors, animal
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (stroma-derived growth factor 1\alpha and 1\beta;
                                                   albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
    Proteins, specific or class
TT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (therapeutic; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Interleukin 1 receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (type 3; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
     Interleukin 1 receptors
ΙT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (type II; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Interferons
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (\alpha ; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
    Chemokine receptors
ΤT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (β chemokine receptor CCR5; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Chemokine receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (β chemokine receptor CCR7; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
    Transforming growth factors
ΤТ
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
```

use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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(\beta 1-; albumin fusion proteins with therapeutic
         proteins for improved shelf-life)
IT
     Transforming growth factors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (\beta 2-; albumin fusion proteins with therapeutic
         proteins for improved shelf-life)
·IT
     Chemokines
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (\beta 9; albumin fusion proteins with therapeutic
         proteins for improved shelf-life)
 ΙT
     Thrombomodulin
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (β; albumin fusion proteins with therapeutic
         proteins for improved shelf-life)
ΙT
     78990-62-2P, Calpain
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (10a and 10b and 10c; albumin fusion proteins with
         therapeutic proteins for improved shelf-life)
     50-56-6P, Oxytocin, biological studies
                                               9002-62-4P, Prolactin, biological
 ΙT
               9002-67-9P, Luteinizing hormone 9002-68-0P, FSH
                                                                    9002-72-6P,
                                                                 9014-42-0P,
                       9004-10-8P, Insulin, biological studies
     Growth hormone
                       11000-17-2P, Vasopressin
                                                11096-26-7P, Erythropoietin
     Thrombopoietin
     33507-63-0P, Substance P
                                 67763-96-6P, Insulin-like growth factor 1
     83869-56-1P, GM-CSF
                           106096-92-8P, Acidic fibroblast growth factor
     106096-93-9P, Basic fibroblast growth factor
                                                     122191-40-6P, ICE
                  123584-45-2P, Fibroblast growth factor 4
                                                              129653-64-1P,
     proteinase
                                   130939-41-2P, Fibroblast growth factor 6
     Fibroblast growth factor 5
     130939-66-1P, Neurotrophin 3 140208-23-7P, Plasminogen activator
                   141760-45-4P, Furin
                                         142243-03-6P, Plasminogen activator
     inhibitor-1
                                         143375-33-1P, Neurotrophin 4
     inhibitor-2
                    143011-72-7P, G-CSF
     148348-14-5P, Fibroblast growth factor 3
                                                 151185-16-9P, Fibroblast growth
                157857-21-1P, Maspin
                                       164003-41-2P, Fibroblast growth factor 8
     factor 9
                                                  187888-07-9P, Endostatin
     185915-22-4P, Fibroblast growth factor 13
     193363-12-1P, Vascular endothelial growth factor D
                                                         203874-76-4P,
                                   204719-95-9P, Fibroblast growth factor 16
     Fibroblast growth factor 12
                                   219563-02-7P, Vascular endothelial growth
     214210-47-6P, Neuropilin 1
                227018-38-4P, Neuropilin 2
                                              271597-10-5P,
     factor E
     Growth/differentiation factor 1
                                        322637-18-3P, Fibroblast growth factor
                                    332350-92-2P, Bone morphogenetic protein
           331718-56-0P, Resistin
     receptor kinase 3
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (albumin fusion proteins with therapeutic proteins
         for improved shelf-life)
 ΙT
     144114-21-6, Retropepsin
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (inhibitors; albumin fusion proteins with
         therapeutic proteins for improved shelf-life)
     127464-60-2P, Vascular endothelial growth factor
 IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
         (isoforms; albumin fusion proteins with therapeutic
         proteins for improved shelf-life)
     127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A
 IT
     protein moiety reduced), full-length or subfragment fusion
     products
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
```

(nucleotide sequence; albumin fusion proteins with

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therapeutic proteins for improved shelf-life)
     155945-98-5, PN: US5962255 SEQID: 59 unclaimed DNA
                                                          156163-00-7
TΤ
     167728-69-0
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     167731-75-1, PN: US5962255 SEQID: 57 unclaimed DNA
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     US5962255 SEQID: 58 unclaimed DNA
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     GenBank A63621
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     367319-60-6
                   367319-66-2
                                 370965-07-4
                                               370965-08-5
     367319-65-1
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
                                 244008-06-8, PN: WO9947540 SEQID: 4 unclaimed
IΤ
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     122024-47-9
           244008-07-9, PN: W09947540 SEQID: 5 unclaimed DNA 244008-08-0, PN:
     WO9947540 SEQID: 6 unclaimed DNA 244008-09-1, PN: WO9947540 SEQID: 7
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     244008-13-7, PN: WO9947540 SEQID: 9 unclaimed DNA 367273-46-9
                  367273-48-1
                                371149-71-2
     367273-47-0
     RL: PRP (Properties)
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     102510-92-9P, Inhibin A
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (\alpha- and \beta-subunits;
                             albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     9061-61-4P, Nerve growth factor
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (β; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    ANSWER 8 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
     2001:781078 HCAPLUS
AN
DN
     135:348850
     Entered STN: 26 Oct 2001
ED
     Albumin fusion proteins with therapeutic proteins for
TI
     improved shelf-life
IN
     Rosen, Craig A.; Haseltine, William A.
PΑ
     Human Genome Sciences, Inc., USA
SO
     PCT Int. Appl., 374 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
IC
     ICM C12N
CC
     63-3 (Pharmaceuticals)
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Section cross-reference(s): 3, 15

والتنفيذ

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                                                           DATE
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
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                    A2
                           20011025
                                          WO 2001-US11924 20010412 ·
    WO 2001079443
PT
    WO 2001079443
                     А3
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            HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
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            RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
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                                         AU 2001-59063
                                                           20010412
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                      Α2
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                                          EP 2001-932546
                                                          20010412
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                          US 2001-833041
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    US 2003125247
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                                                           20010412
    US 2004010134
                      A1
                         20040115
                                          US 2001-833245
                                                           20010412
PRAI US 2000-229358P P
                           20000412
    US 2000-199384P P
                           20000425
    US 2000-256931P P
                           20001221
    WO 2001-US11924
                     W
                           20010412
AB
    The present invention encompasses fusion proteins of
    albumin with various therapeutic proteins. Therapeutic proteins
    may be stabilized to extend the shelf-life, and/or to
    retain the therapeutic protein's activity for extended periods of time in
    solution, in vitro and/or in vivo, by genetically or chemical fusing
    or conjugating the therapeutic protein to albumin or a fragment
    or variant of albumin. Use of albumin fusion
    proteins may also reduce the need to formulate the protein solns. with
    large excesses of carrier proteins to prevent loss of therapeutic proteins
    due to factors such as binding to the container. Nucleic acid mols.
    encoding the albumin fusion proteins of the invention
    are also encompassed by the invention, as are vectors containing these nucleic
    acids, host cells transformed with these nucleic acids vectors, and
    methods of making the albumin fusion proteins of the
     invention and using these nucleic acids, vectors, and/or host cells.
    Thus, plasmid vectors are constructed in which DNA encoding the desired
    therapeutic protein may be inserted for expression of the albumin
    fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA).
    Yeast-derived signal sequences from Saccharomyces cerevisiae invertase
    SUC2 gene, or the stanniocalcin or native human serum albumin
     signal peptides, are used for secretion in yeast or mammalian systems,
    resp. Thus, the fusion product of human growth hormone with
    residues 1-387 of human serum albumin retains essentially intact
    biol. activity after 5 wk of incubation in tissue culture media at
     37°, whereas recombinant human growth hormone used as
     control lost its biol. activity in the first week. Although the potency
    of the albumin fusion proteins is slightly lower than
    the unfused counterparts in rapid bioassays, their biol. stability results
    in much higher biol. activity in the longer term in vitro assay or in vivo
             Addnl., the present invention encompasses pharmaceutical compns.
     comprising albumin fusion proteins and methods of
     treating, preventing, or ameliorating diseases, disorders or conditions
    using albumin fusion proteins of the invention.
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albumin fusion therapeutic protein shelflife

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ΙT
    Bone morphogenetic proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (2; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    Bone morphogenetic proteins
IT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (7; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    Transport proteins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ABC1 (ATP-binding cassette-containing 1); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
    Proteins, specific or class
IT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (ADMP (anti-dorsalizing morphogenetic protein-1); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Agouti signal; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (BPI (bactericidal/permeability-increasing), 21; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
    Transcription factors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (BRCA1; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Transcription factors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (BRCA2; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
    Proteins, specific or class
IT
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (Del-1 (developmentally regulated endothelial locus-1); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (EMAP II (endothelial monocyte activating polypeptide II);
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
ΙT
    Troponins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (I; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Toxins
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)

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Myelin basic protein

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(ML-I (mistletoe lectin I); albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
IT
     Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (MTP (microsomal transfer protein); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (NIF (neutrophil inhibitory factor); albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΤТ
    Receptors
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (T1/ST2; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
    Glycoproteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TNF-BP (tumor necrosis factor-binding protein); albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (TRAIL (tumor necrosis factor-related apoptosis-inducing ligand);
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
IT
     Drug delivery systems
     Gene therapy.
    Molecular cloning
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
IT
    Arrestins
    CD4 (antigen)
    CTLA-4 (antigen)
    Calreticulin
    Cell adhesion molecules
    Ciliary neurotrophic factor
    Cytokines
    Decorins
     Enzymes, biological studies
       Fusion proteins (chimeric proteins)
     Gelsolin
    Growth factors, animal
    Heat-shock proteins
       Interferons
     Interleukin 1
     Interleukin 1 receptor antagonist
     Interleukin 10
     Interleukin 11
     Interleukin 12
     Interleukin 18
     Interleukin 4
     Interleukin 4 receptors
     Interleukin 8
     LFA-3 (antigen)
     Lactoferrins
     Leukemia inhibitory factor
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ΙT

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IT

IT

IT

ΙT

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Platelet-derived growth factors
Pleiotrophins
Stem cell factor
Synthetic gene
Tumor necrosis factor receptors
Tumor necrosis factor receptors
Tumor necrosis factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (albumin fusion proteins with therapeutic proteins
   for improved shelf-life)
Neurotrophic factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (brain-derived; albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (chemokine-binding; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (corticotropin-releasing factor-binding; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (diphtheria, fusion protein with interleukin 2;
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (exotoxins, Pseudomonas, fusion protein with acidic
   fibroblast growth factor; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
Signal peptides
RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
   (for improved secretion in yeast or mammalian cells; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Interleukin 3
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (fusion protein with G-CSF; albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Interleukin 6
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (fusion proteins with diphtheria toxin or Pseudomonas
   exotoxin; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (gene patched; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
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Neurotrophic factors
ΤТ
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (glial-derived; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interferon \omega; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (interferon-induced, 10; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Animal cell
        (mammalian, recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Proteins, specific or class
IT
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (noggins; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
     Plasmid vectors
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Plasmid vectors
        (pScCHSa, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
TΨ
     Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
ΙT
     Hemopoietins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (progenipoietin; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Hemopoietins
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (promegapoietin; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Saccharomyces cerevisiae
ΙT
     Yeast
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
     Antigens
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (retinal S-; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
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Albumins, biological studies

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RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (serum; albumin fusion proteins with therapeutic
  proteins for improved shelf-life)
Genetic element
RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
   (signal sequence, for improved secretion in yeast or mammalian cells;
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
Antibodies
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (single chain; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Hedgehog protein
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (sonic; albumin fusion proteins with therapeutic
  proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (therapeutic; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (tie-2; albumin fusion proteins with therapeutic
  proteins for improved shelf-life)
Complement receptors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (type 1; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Collagens, biological studies
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (type II; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Interferons
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
       albumin fusion proteins with therapeutic
  proteins for improved shelf-life)
Interferons
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (α ; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Transforming growth factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (\beta 1-; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Transforming growth factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (\beta 2-; albumin fusion proteins with therapeutic
   proteins for improved shelf-life)
Transforming growth factors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
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albumin fusion proteins with therapeutic proteins for improved **shelf-life**) IT Interferons RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (γ; albumin fusion proteins with therapeutic proteins for improved **shelf-life**) 139691-92-2P, Serine proteinase inhibitor IT RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (1; albumin fusion proteins with therapeutic proteins for improved shelf-life) 9001-91-6DP, Lys-plasminogen, de-(1-76) derivs. IT RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (Lys-plasminogen; albumin fusion proteins with therapeutic proteins for improved shelf-life) 9002-12-4P, 9001-42-7P, α -Glucosidase 9002-01-1P, Streptokinase ΙT 9002-67-9P, 9002-61-3P, Chorionic gonadotropin Urate oxidase 9002-68-0P, FSH 9002-69-1P, Relaxin 9002-72-6P, Luteinizing hormone Growth hormone 9003-98-9P, DNase 9004-10-8P, Insulin, biological 9007-92-5P, Glucagon, biological studies 9014-42-0P, 9025-35-8P, 9015-68-3P, Asparaginase Thrombopoietin 9026-93-1P, Adenosine deaminase 9035-55-6P, α -Galactosidase 9039-53-6P, Urokinase 9040-61-3P, Staphylokinase 9054-89-1DP, Superoxide dismutase, fusion protein with botulin 9061-61-4P, Nerve growth factor 9073-56-7P, α -L-Iduronidase 11096-26-7P, Erythropoietin 9088-41-9P, Kunitz proteinase inhibitor 37228-64-1P, β -Glucocerebrosidase 42616-25-1P, Methioninase 55354-43-3P, Arylsulfatase B 62229-50-9P, Epidermal growth factor 76901-00-3P, Platelet 67763-96-6P, Insulin-like growth factor 1 activating factor acetylhydrolase 82707-54-8P, Neprilysin 83652-28-2P, Calcitonin gene-related peptide 83869-56-1P, GM-CSF 86090-08-6P, 104625-48-1P, Activin A Angiostatin 99149-95-8P, Saruplase 106096-92-8DP, Acidic 105844-41-5P, Plasminogen activator inhibitor fibroblast growth factor, fusion protein with Pseudomonas 106096-93-9P, Fibroblast growth factor 2 106096-92-8P exotoxin 107231-12-9DP, Botulin, fusion protein with superoxide dismutase 139639-23-9P, 116036-70-5P, Fibrolase 130939-66-1P, Neurotrophin 3 Tissue-type plasminogen activator 143011-72-7P, G-CSF 145137-38-8P, 153858-68-5P, Contortrostatin 157857-21-1P, Maspin Desmoteplase 169494-85-3P, Leptin 186270-49-5P, 163658-39-7P, Prosaptide 194368-66-6P, Angiopoietin 2 194554-71-7P, Tissue Angiopoietin 1 195009-21-3P, Glial growth factor 2 factor pathway inhibitor 197980-93-1P, Pigment epithelium-derived factor 196488-72-9P, Ranpirnase 205944-50-9P, Osteoprotegerin 244019-30-5P, Vascular endothelial growth 320336-96-7P, Kistrin 362605-29-6P, Keratinocyte growth factor 1 factor 1 RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (albumin fusion proteins with therapeutic proteins for improved shelf-life) 9000-95-7P, Apyrase ΙT RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (ecto-; albumin fusion proteins with therapeutic proteins for improved shelf-life) 9002-79-3P, MSH IT RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (fusion products with diphtheria toxin; albumin fusion proteins with therapeutic proteins for improved shelf-life)

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مر والميتوم

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127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A
ΙT
    protein moiety reduced), full-length or subfragment fusion
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (nucleotide sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
                               217893-77-1, GenBank A63614
                                                              217893-78-2,
ΙT
    131748-18-0
                  156163-00-7
    GenBank A63615
                     217893-79-3, GenBank A63616
                                                   217893-80-6, GenBank A63617
                                  217893-82-8, GenBank A63619
    217893-81-7, GenBank A63618
                                                                217893-83-9,
    GenBank A63620
                     217893-84-0, GenBank A63621
                                                  217893-85-1, GenBank A63622
                                217893-89-5, GenBank A63627 217893-90-8,
    217893-86-2, GenBank A63624
                     217893-91-9, GenBank A63629 217893-92-0, GenBank A63630
    GenBank A63628
                  367319-53-7
     367319-52-6
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                                             367319-55-9
                                                            367319-56-0
    367319-58-2
                  367319-59-3
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                                                            367319-62-8
    367319-63-9
                  367319-64-0
                               367319-65-1
                                              367319-66-2
    RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
       proteins with therapeutic proteins for improved shelf-
       life)
    229477-44-5
                  244008-03-5, PN: WO9947540 SEQID: 3 unclaimed DNA
IT
    244008-06-8, PN: WO9947540 SEQID: 4 unclaimed DNA 244008-07-9, PN:
                                       244008-08-0, PN: WO9947540 SEQID: 6
    WO9947540 SEQID: 5 unclaimed DNA
                   244008-09-1, PN: WO9947540 SEQID: 7 unclaimed DNA
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     244008-12-6, 8: PN: WO0183510 SEQID: 8 unclaimed DNA
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    WO9947540 SEQID: 9 unclaimed DNA
                                      244008-14-8, PN: WO9947540 SEQID: 10
    unclaimed DNA
                    367273-46-9
                                 367273-47-0
                                                367273-48-1
                                                            370571-84-9
    RL: PRP (Properties).
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    114949-22-3P, Activin
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (βc; albumin fusion proteins with therapeutic
       proteins for improved shelf-life)
    ANSWER 9 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
ΑN
    2001:781077 HCAPLUS
DN
    135:348849
ED
    Entered STN: 26 Oct 2001
TΙ
    Albumin fusion proteins with therapeutic proteins for
    improved shelf-life
    Rosen, Craig A.; Haseltine, William A.
ΙN
    Human Genome Sciences, Inc., USA
PA
SO
    PCT Int. Appl., 413 pp.
    CODEN: PIXXD2
DT
    Patent
    English
LA
    ICM C12N
IC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 3, 15
FAN.CNT 7
                                          APPLICATION NO.
                                                           DATE
     PATENT NO.
                     KIND DATE
                                          _____
     ______
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                                          WO 2001-US11850 20010412
PΙ
    WO 2001079442
                     Α2
                           20011025
                     A3
                           20020606
    WO 2001079442
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
             VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,

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DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                            20011030
                                           AU · 2001 – 64563
                                                             20010412
    AU 2001064563
                       Α5
                                           EP 2001-938994
     EP 1276849
                       Α2
                            20030122
                                                             20010412
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     US 2003125247
                       Α1
                            20030703
                                           US 2001-833041
                                                             20010412
     US 2003171267
                       A1
                            20030911
                                           US 2001-833117
                                                             20010412
    US 2003199043
                       A1
                            20031023
                                           US 2001-832501
                                                             20010412
                                                             20010412
     JP 2003531590
                       Т2
                            20031028
                                           JP 2001-577426
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                                           US 2001-833118
                                                             20010412
     US 2004010134
                       A1
                            20040115
                                           US 2001-833245
                                                             20010412
PRAI US 2000-229358P
                       Ρ
                            20000412
     US 2000-199384P
                       Ρ
                            20000425
     US 2000-256931P
                       Ρ
                            20001221
     WO 2001-US11850
                       W
                            20010412
     The present invention encompasses fusion proteins of
AB
     albumin with various therapeutic proteins, and in particular
     various antibodies. Therapeutic proteins may be stabilized to extend the
     shelf-life, and/or to retain the therapeutic protein's
     activity for extended periods of time in solution, in vitro and/or in vivo,
     by genetically or chemical fusing or conjugating the therapeutic
     protein to albumin or a fragment or variant of albumin
        Use of albumin fusion proteins may also reduce the
     need to formulate the protein solns. with large excesses of carrier
     proteins to prevent loss of therapeutic proteins due to factors such as
     binding to the container. Nucleic acid mols. encoding the albumin
     fusion proteins of the invention are also encompassed by the
     invention, as are vectors containing these nucleic acids, host cells
     transformed with these nucleic acids vectors, and methods of making the
     albumin fusion proteins of the invention and using these
     nucleic acids, vectors, and/or host cells. Thus, plasmid vectors are
     constructed in which DNA encoding the desired therapeutic protein may be
     inserted for expression of the albumin fusion proteins
     in yeast (pPPC0005) and mammalian cells (pC4:HSA). Yeast-derived signal
     sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the
     stanniocalcin or native human serum albumin signal peptides, are
     used for secretion in yeast or mammalian systems, resp. Thus, the
     fusion product of human growth hormone with residues 1-387 of
     human serum albumin retains essentially intact biol. activity
     after 5 wk of incubation in tissue culture media at 37°, whereas
     recombinant human growth hormone used as control lost its biol.
     activity in the first week. Although the potency of the albumin
     fusion proteins is slightly lower than the unfused counterparts in
     rapid bioassays, their biol. stability results in much higher biol.
     activity in the longer term in vitro assay or in vivo assays. Addnl., the
     present invention encompasses pharmaceutical compns. comprising
     albumin fusion proteins and methods of treating,
     preventing, or ameliorating diseases, disorders or conditions using
     albumin fusion proteins of the invention.
ST
     albumin fusion therapeutic protein shelflife
ΙT
     Antigens
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (17-1A, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (B7.2, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CA125, antibodies to; albumin fusion proteins with
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therapeutic proteins for improved shelf-life)

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ΙT
    CD antigens
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CD147, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CD33, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CD48, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CD52, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (CD6, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
    Immunoglobulins
ΙT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (E, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
    Histocompatibility antigens
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HLA-DR, antibodies to; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HM1.24, antibodies to; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
ΙT
    Cell adhesion molecules
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (ICAM-1 (intercellular adhesion mol. 1), antibodies to; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Immunoglobulin receptors
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (IgG type I, antibodies to; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
ΙT
    Selectins
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (L-, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Integrins
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (LPAM-1 (lymphocyte Peyer's patch high endothelial venule adhesion mol.
        1), antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Blood-group substances
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (Lex, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
    Blood-group substances
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (Ley, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Immunoglobulins
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (M, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
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ΙT

Histocompatibility antigens

ΙT

ΙT

IT

ΙT

ΙT

.IT

ΙT

ΙΤ

IT

ΙΤ

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Angiogenic factors

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RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (MHC (major histocompatibility complex), class I, antibodies to;
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
Histocompatibility antigens
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (MHC (major histocompatibility complex), class II, antibodies to;
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
Proteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (NogoA, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (Nsf2, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Glycoproteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (P170, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Cell adhesion molecules
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (SC-1, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Proteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (SF-25, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Antigens
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (SSEA-1 (stage-specific embryonic antigen 1), antibodies to;
   albumin fusion proteins with therapeutic proteins for
   improved shelf-life)
Antigens
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (TAG-72 (tumor-associated glycoprotein 72), antibodies to; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Cell adhesion molecules
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (VCAM-1, antibodies to; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
Drug delivery systems
Gene therapy
Molecular cloning
   (albumin fusion proteins with therapeutic proteins
   for improved shelf-life)
Antibodies
Cell adhesion molecules
Cytokines
Enzymes, biological studies
  Fusion proteins (chimeric proteins)
Growth factors, animal
Immunoglobulins
  Interferons
Synthetic gene
Tumor necrosis factor receptors
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (albumin fusion proteins with therapeutic proteins
   for improved shelf-life)
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CD14 (antigen)
CD2 (antigen)
CD20 (antigen)
CD22 (antigen)
CD3 (antigen)
CD30 (antigen)
CD38 (antigen)
CD4 (antigen)
CD40 (antigen)
CD44 (antigen)
CD45 (antigen)
CD5 (antigen)
CD8 (antigen)
CD80 (antigen)
CD80 (antigen)
CTLA-4 (antigen)
Carcinoembryonic antigen
Epidermal growth factor receptors
Fas antigen
Integrins
Interleukin 4 receptors
Interleukin 5
LFA-1 (antigen)
Mucins
TCR (T cell receptors)
Transferrin receptors
neu (receptor)
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Mucins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (episialins, antibodies to; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
Signal peptides
RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
   (for improved secretion in yeast or mammalian cells; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Glycoproteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (gD, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Envelope proteins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (gp120env, antibodies to; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
Glycoproteins, specific or class
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (gpII, antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Animal cell
   (mammalian, recombinant expression host; albumin
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Agglutinins and Lectins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (mannan-binding, antibodies to; albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Antibodies
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (monoclonal; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
    Plasmid vectors
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
ΙT
    Plasmid vectors
        (pScCHSa, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
IT
    Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
ΙT
     Interleukin 6 receptors
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (receptor-associated glycoprotein gp130, antibodies to; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΤТ
    Saccharomyces cerevisiae
    Yeast
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Albumins, biological studies
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (serum; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΤT
    Genetic element
    RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (signal sequence, for improved secretion in yeast or mammalian cells;
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
ΙT
    Antibodies
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (single chain; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Venoms
        (snake, antibodies to; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Proteins, specific or class
    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (therapeutic; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Globulins, biological studies
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (thymocyte, antibodies to; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
IT
    Antigens
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (tumor-associated, antibodies to; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
    Interleukin 2 receptors
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ΙT

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unclaimed DNA

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RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\alpha-chain, antibodies to; albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Interferons
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (α ; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
Integrins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\alpha IIb\beta 3, antibodies to;
                            albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Vitronectin receptors
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\alpha v \beta 3, antibodies to;
                           albumin fusion
   proteins with therapeutic proteins for improved shelf-
  ·life)
Integrins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\alpha 4\beta 1, antibodies to;
                           albumin fusion
   proteins with therapeutic proteins for improved shelf-
   life)
Chemokine receptors
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\beta \text{ chemokine receptor CCR5, antibodies to; albumin})
   fusion proteins with therapeutic proteins for improved
   shelf-life)
Integrins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (\beta 2, antibodies to; albumin fusion proteins
   with therapeutic proteins for improved shelf-life)
9002-67-9P, Luteinizing hormone
                                   9002-68-0P, FSH
                                                     9002-72-6P, Growth
          9004-10-8P, Insulin, biological studies
                                                     11096-26-7P,
                 67763-96-6P, Insulin-like growth factor 1
Erythropoietin
                                                             83869-56-1P,
         143011-72-7P, G-CSF
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (albumin fusion proteins with therapeutic proteins
   for improved shelf-life)
156586-89-9
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (albumin fusion proteins with therapeutic proteins
   for improved shelf-life)
                        19600-01-2, Ganglioside GM2
11016-39-0, Properdin
                                                        20830-75-5, Digoxin
99085-47-9, CD55 antigen
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (antibodies to; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A
protein moiety reduced), full-length or subfragment fusion
RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (nucleotide sequence; albumin fusion proteins with
   therapeutic proteins for improved shelf-life)
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                                           167728-72-5
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              167731-74-0, PN: US5962255 SEQID: 56 unclaimed DNA
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US5962255 SEQID: 58 unclaimed DNA
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167731-78-4, PN: US5962255 SEQID: 61 unclaimed DNA

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    RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
       proteins with therapeutic proteins for improved shelf-
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    WO9947540 SEQID: 10 unclaimed DNA
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    367273-48-1
    RL: PRP (Properties)
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
    ANSWER 10 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
    2001:780938 HCAPLUS
    135:322686
    Entered STN: 26 Oct 2001
    Albumin fusion proteins with therapeutic proteins for
    improved shelf-life
    Rosen, Craig A.; Sadeghi, Homayoun; Prior, Christopher P.;
    Turner, Andrew John
    Human Genome Sciences, Inc., USA; Principia Pharmaceutical
    Corporation
    PCT Int. Appl., 328 pp.
    CODEN: PIXXD2
    Patent
    English
    ICM C07K001-00
    ICS A01N037-18
     63-3 (Pharmaceuticals)
    Section cross-reference(s): 3, 15
FAN.CNT 7
                                          APPLICATION NO. DATE
    PATENT NO.
                     KIND DATE
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                                          WO 2001-US12008 20010412
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            CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
            HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
            LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
            RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
            VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
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BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

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                                           EP 2001-932549
     EP 1274720
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                            20030703
                                                             20010412 .
     US 2003125247
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                            20030911
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                                                             20010412
     US 2003171267
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                       T2
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                                                             20010412
                            20031023
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                                           US 2001-833118
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                                                             20010412
                                           US 2001-833245
     US 2004010134
                       Α1
                            20040115
                                                             20010412
PRAI US 2000-229358P
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     US 2000-199384P
                       Ρ
                            20000425
     US 2000-256931P
                       Ρ
                            20001221
    WO 2001-US12008 W
                            20010412
     The present invention encompasses fusion proteins of
AB
     albumin with various therapeutic proteins, and in particular, with
     interleukin 2, calcitonin, growth hormone-releasing factor,
     interferon \boldsymbol{\beta} , parathyroid hormine, and insulin-like
     growth factor 1. Therapeutic proteins may be stabilized to extend the
     shelf-life, and/or to retain the therapeutic protein's
     activity for extended periods of time in solution, in vitro and/or in vivo,
     by genetically or chemical fusing or conjugating the therapeutic
     protein to albumin or a fragment or variant of albumin
        Use of albumin fusion proteins may also reduce the
     need to formulate the protein solns. with large excesses of carrier
    proteins to prevent loss of therapeutic proteins due to factors such as
     binding to the container. Nucleic acid mols. encoding the albumin
     fusion proteins of the invention are also encompassed by the
     invention, as are vectors containing these nucleic acids, host cells
     transformed with these nucleic acids vectors, and methods of making the
     albumin fusion proteins of the invention and using these
     nucleic acids, vectors, and/or host cells. Thus, plasmid vectors are
     constructed in which DNA encoding the desired therapeutic protein may be
     inserted for expression of the albumin fusion proteins
     in yeast (pPPC0005) and mammalian cells (pC4:HSA).
                                                          Yeast-derived signal
     sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the
     stanniocalcin or native human serum albumin signal peptides, are
     used for secretion in yeast or mammalian systems, resp. Thus, the
     fusion product of human growth hormone with residues 1-387 of
     human serum albumin retains essentially intact biol. activity
     after 5 wk of incubation in tissue culture media at 37°, whereas
     recombinant human growth hormone used as control lost its biol.
     activity in the first week. Although the potency of the albumin
     fusion proteins is slightly lower than the unfused counterparts in
     rapid bioassays, their biol. stability results in much higher biol.
     activity in the longer term in vitro assay or in vivo assays. Addnl., the
     present invention encompasses pharmaceutical compns. comprising
     albumin fusion proteins and methods of treating,
     preventing, or ameliorating diseases, disorders or conditions using
     albumin fusion proteins of the invention.
ST
     albumin fusion therapeutic protein shelflife
ΙT
     Hepatitis
        (C, agents for treatment of; albumin fusion
        proteins with therapeutic proteins for improved shelf-
ΙT
     Antitumor agents
        (Kaposi's sarcoma; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
     Antitumor agents
IT
        (acute myelogenous leukemia; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
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والمتناث

Anti-AIDS agents

Antidiabetic agents

ΙT

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```
Antirheumatic agents
     Drug delivery systems
    Gene therapy
     Immunosuppressants
    Molecular cloning
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
TΤ
    Cell adhesion molecules
    Cytokines
    Enzymes, biological studies
      Fusion proteins (chimeric proteins)
    Growth factors, animal
       Interferons
     Interleukin 2
     Synthetic gene
     Tumor necrosis factor receptors
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
     Signal peptides
IT
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (for improved secretion in yeast or mammalian cells; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
IT
     Intestine, disease
        (inflammatory, agents for treatment of; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
ΙT
    Kidney, neoplasm
     Lung, neoplasm
     Ovary, neoplasm
        (inhibitors; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
    Antitumor agents
        (kidney; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Antitumor agents
        (leukemia; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Antitumor agents
        (lung; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Animal cell
        (mammalian, recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Antitumor agents
ΙT
        (melanoma, metastasis; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Antitumor agents
        (melanoma; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
ΙT
     Antitumor agents
        (non-Hodgkin's lymphoma; albumin fusion proteins
        with therapeutic proteins for improved shelf-life)
IT
     Antitumor agents
        (ovary; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
     Plasmid vectors
        (pC4:HSA, for mammalian cell expression; albumin
```

fusion proteins with therapeutic proteins for improved

```
shelf-life)
IT
    Plasmid vectors
        (pPPC0005, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
    Plasmid vectors
IT
        (pScCHSa, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
     Plasmid vectors
        (pScNHSA, for yeast expression; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     Saccharomyces cerevisiae
ΙT
     Yeast
        (recombinant expression host; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
     Kidney, neoplasm
IT
        (renal-cell carcinoma, metastasis, inhibitors; albumin
        fusion proteins with therapeutic proteins for improved
        shelf-life)
TT
    Antitumor agents
        (renal-cell carcinoma, metastasis; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
IT
    Albumins, biological studies
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (serum; albumin fusion proteins with therapeutic
        proteins for improved shelf-life)
IT
    Genetic element
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (signal sequence, for improved secretion in yeast or mammalian cells;
        albumin fusion proteins with therapeutic proteins for
        improved shelf-life)
IT
    Antibodies
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (single chain; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
IT
    Multiple sclerosis
        (therapeutic agents; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Proteins, specific or class
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (therapeutic; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (\alpha ; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (\beta ; albumin fusion proteins with
       therapeutic proteins for improved shelf-life)
     9002-64-6P, Parathyroid hormone 9002-67-9P, Luteinizing hormone
ΙT
     9002-68-0P, FSH
                     9002-72-6P, Growth hormone
                                                    9004-10-8P, Insulin,
```

9007-12-9P, Calcitonin

9034-39-3P, Growth

biological studies

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و المستقبل

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CC

63-3 (Pharmaceuticals)

```
11096-26-7P, Erythropoietin
     hormone-releasing factor
                                                               67763-96-6P,
     Insulin-like growth factor 1
                                   83869-56-1P, GM-CSF
                                                          143011-72-7P, G-CSF
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins
        for improved shelf-life)
     127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A
     protein moiety reduced), full-length or subfragment fusion
     RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (nucleotide sequence; albumin fusion proteins with
       therapeutic proteins for improved shelf-life)
ΙT
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     GenBank A63618
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     244008-14-8, PN: WO9947540 SEQID: 10 unclaimed DNA
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     RL: PRP (Properties)
        (unclaimed nucleotide sequence; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     367510-76-7
ΙT
     RL: PRP (Properties)
        (unclaimed protein sequence; albumin fusion
        proteins with therapeutic proteins for improved shelf-
        life)
     131748-18-0
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IT
                   367273-46-9
     RL: PRP (Properties)
        (unclaimed sequence; albumin fusion proteins with
        therapeutic proteins for improved shelf-life)
              THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Beth Israel Hospital Association; WO 9618412 Al 1996 HCAPLUS
(2) Lee; Pharm Dev Tech 1999, V4(2), P269 HCAPLUS
(3) Rhone-Poulenc Rorer S A; WO 9315199 A1 1993 HCAPLUS
(4) Rhone-Poulenc Rorer S A; WO 9315211 A1 1993 HCAPLUS
(5) Takahashi; Peptides 1997, V18(3), P439 HCAPLUS
(6) Yeh; Prc Nat Acad Sci USA 1992, V69, P1904
    ANSWER 11 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
ΑN
     2001:763025
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DN
     135:335111
ED
     Entered STN: 19 Oct 2001
     Albumin fusion proteins with therapeutic proteins for improved shelf-life
ΤI
IN
     Rosen, Craig A.; Haseltine, William A.
PA
     Human Genome Sciences, Inc., USA
SO
     PCT Int. Appl., 2102 pp.
     CODEN: PIXXD2
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     Patent
     English
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IC
     ICM C07H021-04
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Section cross-reference(s): 3, 15

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                     A1 20040115
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PRAI US 2000-229358P P
                          20000412
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     US 2000-199384P
                           20000425
                    Р
                           20001221
     US 2000-256931P
     WO 2001-US11988
                    W
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The present invention encompasses fusion proteins of albumin with various therapeutic proteins. Therapeutic proteins may be stabilized to extend the shelf-life, and/or to retain the therapeutic protein's activity for extended periods of time in solution, in vitro and/or in vivo, by genetically or chemical fusing or conjugating the therapeutic protein to albumin or a fragment or variant of albumin. Use of albumin fusion proteins may also reduce the need to formulate the protein solns. with large excesses of carrier proteins to prevent loss of therapeutic proteins due to factors such as binding to the container. Nucleic acid mols. encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Thus, plasmid vectors are constructed in which DNA encoding the desired therapeutic protein may be inserted for expression of the albumin fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA). Yeast-derived signal sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the stanniocalcin or native human serum albumin signal peptides, are used for secretion in yeast or mammalian systems, resp. Thus, the fusion product of human growth hormone with residues 1-387 of human serum albumin retains essentially intact biol. activity after 5 wk of incubation in tissue culture media at 37°, whereas recombinant human growth hormone used as control lost its biol. activity in the first week. Although the potency of the albumin fusion proteins is slightly lower than the unfused counterparts in rapid bioassays, their biol. stability results in much higher biol. activity in the longer term in vitro assay or in vivo assays. Addnl., the present invention encompasses pharmaceutical compns. comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention.

ST albumin fusion therapeutic protein shelflife

IT Drug delivery systems

Gene therapy

Molecular cloning

(albumin fusion proteins with therapeutic proteins for improved shelf-life)

IT Cell adhesion molecules

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- F. .

IT

improved shelf-life)
9002-67-9P, Luteinizing hormone

Cytokines Enzymes, biological studies Fusion proteins (chimeric proteins) Growth factors, animal Interferons Synthetic gene Tumor necrosis factor receptors RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT Signal peptides RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (for improved secretion in yeast or mammalian cells; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT (mammalian, recombinant expression host; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT Plasmid vectors (pC4:HSA, for mammalian cell expression; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT (pPPC0005, for yeast expression; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT Plasmid vectors (pScCHSa, for yeast expression; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT (pScNHSA, for yeast expression; albumin fusion proteins with therapeutic proteins for improved shelf-life) Saccharomyces cerevisiae IT Yeast (recombinant expression host; albumin fusion proteins with therapeutic proteins for improved shelf-life) Albumins, biological studies IT RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (serum; albumin fusion proteins with therapeutic proteins for improved shelf-life) Genetic element TT : RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (signal sequence, for improved secretion in yeast or mammalian cells; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT Antibodies RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (single chain; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙΤ Proteins, specific or class RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (therapeutic; albumin fusion proteins with therapeutic proteins for improved shelf-life) ΙT Interferons RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) $(\alpha;$ albumin fusion proteins with therapeutic proteins for

9002-68-0P, FSH

9004-10-8P, Insulin, biological studies

9002-72-6P, Growth

11096-26-7P,

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67763-96-6P, Insulin-like growth factor 1
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             143011-72-7P, G-CSF
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    use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (albumin fusion proteins with therapeutic proteins for improved
        shelf-life)
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     127361-02-8DP, Albumin (human blood serum clone HSA-II/HSA-I-A protein
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    RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic
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        (unclaimed nucleotide sequence; albumin fusion proteins with
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        proteins for improved shelf-life)
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369638-84-6

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(unclaimed sequence; albumin fusion proteins with therapeutic proteins for improved shelf-life)

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THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 3
(1) Delta Biotechnology Limited; EP 0322094 A1 1989 HCAPLUS
(2) Delta Biotechnology Limited; WO 9724445 A1 1997 HCAPLUS
(3) Human Genome Sciences Inc; WO 9734997 A1 1997 HCAPLUS
    ANSWER 12 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
ΑN
     2000:609058 HCAPLUS
DN
     133:168425
     Entered STN: 01 Sep 2000
ED
     Suppository of recombinant human interferon .
ΤI
     alpha.2a
     Chen, Weijia; Zheng, Hui; Zhang, Yan; Wang, Dongqian
IN
     Changchun Biological Product Inst., Ministry of Public Health, Peop. Rep.
PΑ
SO
     Faming Zhuanli Shenqing Gongkai Shuomingshu, 5 pp.
     CODEN: CNXXEV
DT
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     Chinese
     ICM A61K009-02
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     63-6 (Pharmaceuticals)
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PRAI CN 1999-105589
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     Suppository of interferon \alpha 2a comprise
     recombinant human interferon \alpha 2a solution
     (0.5 MIU per suppository) 14, glycerol 58, gelatin 26, and human serum
     albumin 2%. The preparation process involves mixing glycerol with
     gelatin, standing overnight, sterilizing for 20-30 min, cooling to
     40-56\Phi', adding recombinant human interferon .
     alpha.2a, and shaping.
     recombinant human interferon alpha 2a
ST
     suppository
IT
     Albumins, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (serum; suppository of recombinant human interferon
        \alpha 2a)
ΙT
     Drug delivery systems
        (suppositories; suppository of recombinant human
        interferon \alpha 2a)
ΙT
     Anti-inflammatory agents
     Antitumor agents
     Antiviral agents
     Skin, disease
        (suppository of recombinant human interferon
        \alpha 2a)
IT
     Gelatins, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (suppository of recombinant human interferon
        \alpha 2a)
ΙT
     Interferons
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        (α -2a, recombinant human;
        suppository of recombinant human interferon
        \alpha 2a)
     56-81-5, Glycerol, biological studies
IT
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        (suppository of recombinant human interferon
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ور وأنتقاء

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    ANSWER 13 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
     1999:783954 HCAPLUS
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ED
     Entered STN: 10 Dec 1999
     Recombinant human interferon \beta -1A (
TI
     IFN-beta-1A) formulation
IN
     Alam, John; Rogge, Mark; Goelz, Susan
    Biogen, Inc., USA
PΑ
SO
     PCT Int. Appl., 28 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
IC
     ICM A61K038-21
     63-6 (Pharmaceuticals)
     Section cross-reference(s): 15
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             UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
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     Liquid compns. comprising a buffer of pH about 7.2, recombinant
     interferon-\beta and 15 mg/mL of human serum
     albumin, and kits for parenteral administration comprising said
     compns. are disclosed.
     recombinant interferon beta formulation
ST
IT
     Medical goods
        (alc. swabs; recombinant human interferon
        \beta -1A (IFN-beta-1A) formulation)
ΙT
     Medical goods
        (bandages, adhesive; recombinant human interferon
        \beta -1A (IFN-beta-1A) formulation)
ΙT
     Buffers
     Molecular cloning
     Needles (tools)
     Syringes
     рΗ
        (recombinant human interferon \beta -1A (
        IFN-beta-1A) formulation)
     Albumins, biological studies
IΤ
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL: (Biological study); PROC (Process); USES (Uses)
        (serum, human; recombinant human interferon
        \beta -1A (IFN-beta-1A) formulation)
ΙT
     Interferons
     RL: BPN (Biosynthetic preparation); PEP (Physical, engineering or chemical
     process); THU (Therapeutic use); BIOL (Biological study); PREP
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(Preparation); PROC (Process); USES (Uses)
        (β ; recombinant human interferon
        β -1A (IFN-beta-1A) formulation)
     145258-61-3, Interferon \beta 1 (human fibroblast
IT
     protein moiety)
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL (Biological study); PROC (Process); USES (Uses)
        (recombinant human interferon β -1A (
        IFN-beta-1A) formulation)
              THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Alam, J; Pharmaceutical Research 1997, V14(4), P546 HCAPLUS
(2) Anon; http://www.healthdirect.com/usenew/pressrel/p biogel.htm 1996
(3) Salmon, P; Journal of Interferon and Cytokine Research 1996, V16(10), P759
   HCAPLUS
(4) US Food and Drug Administration-Interferon Beta-1A, Biogen, Inc;
   http://www.fda.gov/cber/products/ifnbbio051796.htm,
   http://www.fda.gov/cber/label/infbbio051796lb.pdf 1998
L66 ANSWER 14 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
     1999:563880 HCAPLUS
ΑN
DN
     131:161626
                   08 Sep 1999
ED
     Entered STN:
ΤI
     Oral recombinant human \alpha -interferon
     compositions
     Dong, Yilan; Cheng, Xiaogeng; Lin, Yuxin; Wang, Shiwen; Liu, Zhenhao;
IN
PA
     Changchun Institute of Biological Products, Ministry of Public Health,
     Peop. Rep. China
     Faming Zhuanli Shenging Gongkai Shuomingshu, 8 pp.
SO
     CODEN: CNXXEV
DΤ
     Patent
LA
     Chinese
IC
     ICM A61K038-21
CC
     63-6 (Pharmaceuticals)
     Section cross-reference(s): 1, 15
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                             DATE
     ______
                      ____
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                                            ______
     CN 1116951
                            19960221
                                           CN 1995-101216
                                                             19950125 <--
                       Α
PRAI CN 1995-101216
                            19950125
                                      <--
     Title compns. as antiviral agents contain recombinant human .
     alpha.-interferon 100-500 IU, thymosin F5 isolated from
     calf's thymus gland 1-20 \mu g, stabilizers and conventional medical
     additives. The stabilizers are selected from human serum albumin
     , cattle serum albumin, \beta-cyclodextrin and PEG 800.
ST
     recombinant human interferon tablet antiviral
ΙT
     Antiviral agents
     Stabilizing agents
        (oral recombinant human \alpha -interferon
        compns.)
ΙT
     Polyoxyalkylenes, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (oral recombinant human α -interferon
        compns.)
ΙT
     Drug delivery systems
        (oral; oral recombinant human \alpha -
        interferon compns.)
     Albumins, biological studies
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (serum, human or bovine; oral recombinant human
        \alpha -interferon compns.)
IT
     Drug delivery systems
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F. ..

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(tablets; oral recombinant human \alpha -
        interferon compns.)
IT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (α , recombinant human; oral
        recombinant human α -interferon
        compns.)
ΙT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
       . (α -2a, recombinant human; oral
        recombinant human α -interferon
        compns.)
TΤ
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (α -2b, recombinant human; oral
        recombinant human α -interferon
        compns.)
ΙT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (α 1, recombinant human; oral
        recombinant human α -interferon
        compns.)
IT
     61512-21-8, Thymosin
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (F5; oral recombinant human \alpha -
        interferon compns.)
     7585-39-9, \beta-Cyclodextrin
                                 25322-68-3
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (oral recombinant human \alpha -interferon
        compns.)
    ANSWER 15 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
     1997:756962 HCAPLUS
AN
DN
     128:16442
     Entered STN: 04 Dec 1997
ED
     Stabilization of interferons in aqueous solution for manufacture
TI
     of sublingually administered tablets
     Rothschild, Peter R.
IN
PΑ
     Feronpatent Limited, Ire.; Rothschild, Peter R.
SO
     PCT Int. Appl., 12 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
     ICM A61K038-21
IC
     ICS A61K009-20
     63-6 (Pharmaceuticals)
FAN.CNT 1
                      KIND DATE
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                                           -----
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                           _____
                                                            _____
                                                            19970509 <--
                            19971113
                                           WO 1997-IB531
PΤ
     WO 9741885
                       A1
             AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ,
             LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL,
             PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ,
             VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB,
             GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN,
             ML, MR, NE, SN, TD, TG
                                           AU 1997-24011
                                                             19970509 <--
     AU 9724011
                            19971126
                       Α1
                                           EP 1997-919596
                                                             19970509 <--
     EP 920329
                            19990609
                       Α1
                            20020925
     EP 920329
                       В1
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
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E
                                           AT 1997-919596
                                                             19970509 <--
    AT 224725
                            20021015
     ES 2184084
                       Т3
                            20030401
                                           ES 1997-919596
                                                             19970509 <--
                            19960509
PRAI WO 1996-IB433
                       Α
                                     <--
     WO 1997-IB531
                       W
                            19970509
                                      <--
AB
    Natural and recombinant interferons are stabilized
    with bidistd. water, lactose, albumin, sodium mono- and
     dihydrogen phosphates, (C5H10O5)n, such as arabic gum, dissolved and diluted
     in 20 % ethanol solution to the fourth decimal by homeopathic method. The
     final solution is sprayed on to an excipient comprising of 20 % arabic gum,
     30 % lactose and 50 % starch for manufacturing tablets of 100 mg each
containing 200
     I.U. of human alfa-interferon. The tablets are sublingually
     administered to the patient for treatment of viral infections
     sensitive to interferon. Preparation of sublingual tablets according
     above method is disclosed.
     stabilization interferon polysaccharide sublingual
ST
     pharmaceutical tablet
IT
     Hepatitis
        (B; stabilization of interferons in aqueous solution for manufacture of
        sublingually administered tablets)
ΙT
        (C; stabilization of interferons in aqueous solution for manufacture of
        sublingually administered tablets)
IT
        (homeopathy; stabilization of interferons in aqueous solution for
        manufacture of sublingually administered tablets)
TΤ
     Antitumor agents
     Stabilizing agents
        (stabilization of interferons in aqueous solution for manufacture of
        sublingually administered tablets)
    Albumins, biological studies
TΤ
       Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stabilization of interferons in aqueous solution for manufacture of
        sublingually administered tablets)
     Drug delivery systems
IT
        (tablets, sublingual; stabilization of interferons in aqueous
        solution for manufacture of sublingually administered tablets)
ΙT
        (viral; stabilization of interferons in aqueous solution for manufacture
        of sublingually administered tablets)
IT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\alpha ; stabilization of interferons in aqueous solution
        for manufacture of sublingually administered tablets)
ΙT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\beta ; stabilization of interferons in aqueous solution
        for manufacture of sublingually administered tablets)
IT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\gamma; stabilization of interferons in aqueous solution for
        manufacture of sublingually administered tablets)
     63-42-3, Lactose
                       7558-79-4, Sodium monohydrogen phosphate
                                                                    7558-80-7,
IT
     Sodium dihydrogen phosphate 9000-01-5, Arabic gum
                                                           9005-25-8, Starch,
     biological studies
    RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stabilization of interferons in aqueous solution for manufacture of
        sublingually administered tablets)
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L66 ANSWER 16 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN

AN **1996:635884** HCAPLUS

DN 125:308823

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JP 10500125

T2

19980106

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Entered STN: 28 Oct 1996
ΕD
TТ
     Shelf-life of recombinant human interferon .
     alpha.2b under different storage conditions
     Barberia, Daisy; Vega, Maribel; Ferrero, Joel; Duany, Lady; Moya, Galina;
ΑU
     Curras, Tania; Martinez, Maida; Cruz, Asterio; Gil, Miriela; Quintana,
     Marisel
     Centro de Ingenieria Genetica y Biotecnologia, Havana, Cuba
CS
     Biotecnologia Aplicada (1996), 13(3), 190-194
SO
     CODEN: BTAPEP; ISSN: 0864-4551
PΒ
     Sociedad Iberolatinoamericana de Biotecnologia Aplicada a la Salud
DT
     Journal
LA
     Spanish
CC.
     63-5 (Pharmaceuticals)
AB
     The stability test studies under accelerated and normal storage conditions
     carried out with recombinant human alpha 2b interferon
     (hu-r alpha 2b IFN) in phosphate buffer 0.1M, pH 7.0, with and without
     albumin, in order to establish its shelf-life at refrigerating and
     frozen conditions. According to the accelerated study the authors
     concluded that no alterations will interfere with the recognition of hu-r
     alpha 2b IFN in ELISA in at least five years when stored at -70 or
     -20°. Otherwise, when stored at 4°, a loss of 10% may occur
     in one year. The authors corroborated this when the presence of new
     structures which might affect the protein immunol. recognition were
     detected by RP-HPLC. No stabilizing properties of albumin on
     hu-r alpha 2b IFN were observed at least when it is in phosphate buffer 0.1M,
     pH 7.0 and under accelerated storing conditions.
ST
     interferon stability denaturation freezing
ΙT
     Albumins, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (shelf-life of recombinant human interferon
        α 2b under different storage conditions)
ΙT
     Interferons
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\alpha - 2b, shelf-life of recombinant
        human interferon α 2b under
        different storage conditions)
    ANSWER 17 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
ΑN
     1996:43019 HCAPLUS
DN
     124:66661
     Entered STN: 23 Jan 1996
ED
     Stabilized \beta -interferon liquid formulations
ΤI
ΙN
     Samaritani, Fabrizio; Natale, Patrizia
PA
     Applied Research Systems ARS Holding N.V., Neth.
SO
     PCT Int. Appl., 17 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
     ICM A61K038-21
IC
CC
     63-6 (Pharmaceuticals)
FAN.CNT 1
     PATENT NO.
                                           APPLICATION NO.
                                                             DATE
                      KIND DATE
     ______
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                                           _____
     WO 9531213
                      Α1
                            19951123
                                           WO 1995-EP1825
                                                             19950515 <--
         W: AU, CA, JP, US
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
     CA 2190465
                       AA
                            19951123
                                           CA 1995-2190465 19950515 <--
     AU 9526704
                            19951205
                                           AU 1995-26704
                                                             19950515 <--
                       Α1
     AU 704827
                       B2
                            19990506
     EP 759775
                       Α1
                            19970305
                                           EP 1995-921749
                                                             19950515 <--
                       В1
                            20000726
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE

JP 1995-529360

19950515 <--

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AT 194917
                            2000081.5
                                            AT 1995-921749
                                                             19950515 <--
                       Т3
                                            ES 1995-921749
                                                             19950515 <--
     ES 2148526
                            20001016
                            19940516
PRAI IT 1994-RM300
                       Α
                                      <--
    WO 1995-EP1825
                       W
                            19950515
                                      <--
     \beta -Interferon liquid formulations are stabilized
AB
     with a polyol, a nonreducing sugar, or an amino acid. In particular, the
     formulations are stabilized with a polyol, such as mannitol. The
     formulations, preferably, furthermore comprise a buffer, such as acetate
     buffer at a pH 3-4 and human albumin at a min. quantity. The .
    beta.-interferon is preferably recombinant.
     interferon soln stabilizer polyol albumin buffer;
ST
    mannitol albumin acetate buffer interferon stability
IT
     Buffer substances and systems
        (acetate; stabilized \beta -interferon liquid
        formulations)
ΙT
     Albumins, biological studies
     Amino acids, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stabilized \beta -interferon liquid formulations)
     Carbohydrates and Sugars, biological studies
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (nonreducing, stabilized \beta -interferon liquid
        formulations)
     Alcohols, biological studies
IT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (polyhydric, stabilized \beta -interferon liquid
        formulations)
ΙT
     Pharmaceutical dosage forms
        (solns., stabilized \beta -interferon liquid
        formulations)
     Interferons
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (\beta , recombinant; stabilized \beta -
        interferon liquid formulations)
     56-40-6, Glycine, biological studies
                                            57-50-1, Saccharose, biological
ΙT
             69-65-8, D-Mannitol
     studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stabilized \beta -interferon liquid formulations)
    ANSWER 18 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
1.66
AN
     1995:498838 HCAPLUS
DN
     122:248213
     Entered STN: 20 Apr 1995
ΕD
     Influence of human serum albumin content in
TΤ
     formulations on the bioequivalency of interferon alfa-2a given
     by subcutaneous injection in healthy male volunteers
     Zhi, Jianquo; Teller, Stuart B.; Satoh, Hiroko; Koss-Twardy, Susan G.;
ΑU
     Luke, David R.
     Department of Clinical Pharmacokinetics, Hoffmann-La Roche, Inc., Nutley,
CS
     NJ, 07110-1199, USA
     Journal of Clinical Pharmacology (1995), 35(3), 281-4
SO
     CODEN: JCPCBR; ISSN: 0091-2700
DT
     Journal
     English
LΑ
     63-6 (Pharmaceuticals)
CC
     Section cross-reference(s): 1
     To determine the influence of human serum albumin (HSA)
AΒ
     content in formulations on the bioequivalency of recombinant
     interferon \alpha 2a, a double-blind, randomized,
     two-way crossover study was conducted in 24 healthy male volunteers.
     Subjects received a single s.c. injection of 18 million IU of Roferon-A
     reconstituted with either the diluent containing 10 mg of HSA or the HSA-free
     diluent; final HSA contents in the 2 formulations were 15 and 5 mg, resp.
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Administration of the 2 formulations resulted in similar 48-h Roferon-A serum concentration-time profiles and comparable frequency and intensity of adverse events. The statistical anal. using the two one-sided tests procedure showed that both formulations were bioequivalent for pharmacokinetic parameters such as Cmax, tmax, AUC48, and AUC. threefold change in HSA content in formulations does not alter the bioequivalency of Roferon-A. interferon bioavailability bioequivalence injection albumin Drug bioavailability (human serum albumin effect on bioequivalence of recombinant interferon α 2a from s.c. injection in humans) Albumins, biological studies RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (human serum albumin effect on bioequivalence of recombinant interferon α 2a from s.c. injection in humans) Pharmaceutical dosage forms (injections, s.c., human serum albumin effect on bioequivalence of recombinant interferon α 2a from s.c. injection in humans) Interferons RL: BPR (Biological process); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) $(\alpha$ -2a, human serum albumin effect on bioequivalence of recombinant interferon α 2a from s.c. injection in humans) L66 ANSWER 19 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN **1994:6892** HCAPLUS 120:6892 Entered STN: 08 Jan 1994 Novel recombinant human IFN- β , its preparation, and pharmaceutical compositions containing it Siklosi, Thomas; Joester, Karl-eduard; Hofer, Hans BIOFERON Biochemische Substanzen GmbH und Co, Germany Eur. Pat. Appl., 19 pp. CODEN: EPXXDW Patent German ICM C07K015-26 ICS C07K003-28; A61K037-66 16-2 (Fermentation and Bioindustrial Chemistry) Section cross-reference(s): 15 FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE -----______ ----EP 529300 A1 19930303 EP 1992-112427 19920721 <--EP 529300 B1 19981014 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, MC, NL, PT, SE DE 4128319 A1 19930304 DE 1991-4128319 19910827 <--AT 172206 19981015 AT 1992-112427 19920721 <--E. ES 2121804 Т3 19981216 ES 1992-112427 19920721 <--PRAI DE 1991-4128319 19910827 <--A recombinant human β -interferon (IFN- β) produced in mammalian cells, whose oligosaccharide component comprises biantennary ≥60%, triantennary ≥15%, and tetraantennary 0-5% and contains fucose and ≥80% sialic acid, is useful for treatment of tumors, especially Kaposi's sarcoma. Thus, recombinant IFN- β was produced in transfected CHO BIC 8622 cells in MEM containing fetal calf serum and secreted

into the medium in a yield of 1 + 105-1 + 106 IU/L. The

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ΙT

ΙT

ΙT

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IFN-β was purified by liquid-liquid extraction in a PEG
2000-salt solution system, affinity chromatog. on Blue Dextran FF, metal
chelate chromatog. on a Zn2+-loaded chelating Sepharose column, and size
exclusion chromatog. on Sephacryl. The product showed a purity of >99%
and high stability at -20, +15, or +25° when mixed with buffered
human serum albumin and stored for 1-4 wk. Enzymic removal of
terminal sialic acid residues diminished the stability.
recombinant beta interferon purifn
Polyoxyalkylenes, biological studies
Salts, biological studies
RL: BIOL (Biological study)
   (in \beta -interferon purification, by partition)
Oligosaccharides
Sialic acids
RL: BIOL (Biological study)
   (of recombinant \beta -interferon)
Chromatography, gel
   (of β -interferon)
Partition
   (of \beta -interferon, in polyalkylene
   glycol/dextran and polyalkylene glycol/salt systems)
Neoplasm inhibitors
   (recombinant \beta - interferon)
Dyes
   (\beta -interferon affinity chromatog. on)
Animal cell line
   (CHO, recombinant β -interferon
   manufacture with)
Neoplasm inhibitors
   (Kaposi's sarcoma, recombinant \beta -
   interferon as)
Chromatography, column and liquid
   (affinity, of \beta -interferon, on dye)
Coordination compounds
RL: BIOL (Biological study)
   (chelates, stationary phases containing, for \beta -
   interferon chromatog.)
Interferons
RL: BIOL (Biological study)
   (\beta , purification of recombinant, for Kaposi's
   sarcoma treatment)
                                                57-55-6, 1,2-Propanediol,
             148498-83-3, Blue Sepharose FF
12236-82-7
       107-21-1, 1,2-Ethanediol, uses
RL: BIOL (Biological study)
   (in \beta -interferon purification, by affinity
   chromatog.)
                          71-00-1, Histidine, uses
                                                       288-32-4, Imidazole, ·
56-40-6, Glycine, uses
RL: USES (Uses)
   (in \beta -interferon purification, by metal chelate
   chromatog.)
                           68-04-2, Sodium citrate
                                                       25322-68-3,
62-76-0, Sodium oxalate
Polyethylene glycol 25322-69-4, Polypropylene glycol 7447-40-7, Potassium chloride (KCl), uses 7447-41-8, Lithium chloride, uses
                                                            7447-40-7,
                                 7558-80-7, Sodium dihydrogen phosphate
7558-79-4, Disodium phosphate
7647-14-5, Sodium chloride, uses 7681-11-0, Potassium iodide, uses
                                  7757-82-6, Sodium sulfate, uses
7681-82-5, Sodium iodide, uses
7758-11-4, Dipotassium phosphate 7778-80-5, Potassium sulfate, uses
7783-20-2, Ammonium sulfate, uses 9004-54-0, Dextran, uses
                                                                 12125-02-9,
Ammonium chloride, uses
RL: BIOL (Biological study)
   (in \beta -interferon purification, by partition)
131-48-6, N-Acetylneuraminic acid 1113-83-3 2438-80-4, Fucose
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83412-55-9
     32181-59-2, N-Acetyllactosamine
                                       78392-81-1
                                                                  84813-89-8
                   131432-29-6 148553-76-8 148553-77-9 148553-78-0
     123618-73-5
                                 148553-81-5
     148553-79-1
                   148553-80-4
                                               148614-65-7
                                                              148615-15-0
     RL: BIOL (Biological study)
        (of recombinant \beta -interferon)
     7440-02-0D, Nickel, chelates 7440-48-4D, Cobalt, chelates
ΙT
                                                                    7440-50-8D,
     Copper, chelates
                       7440-66-6D, Zinc, chelates 12774-36-6, Sephadex G150
     97599-42-3, Superose 12
                               119332-87-5, Sephacryl S 200 High Resolution
     148499-25-6, TSK-SW 3000
     RL: BIOL (Biological study)
        (\beta -interferon purification by chromatog. on)
     ANSWER 20 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
     1992:468225 HCAPLUS
ΑN
DN
     117:68225
ED
     Entered STN:
                  23 Aug 1992
TΙ
     Human \beta -interferon incubated with muscle
     homogenate is protected by albumin but not by proteinase
ΑU
     Paulesu, L.; Pessina, G. P.; Bocci, V.
     Inst. Gen. Physiol., Univ. Siena, Siena, 53100, Italy
CS
     Proceedings of the Society for Experimental Biology and Medicine (
SO
     1992), 200(3), 414-17
     CODEN: PSEBAA; ISSN: 0037-9727
DT
     Journal
LA
     English
CC
     15-5 (Immunochemistry)
     Section cross-reference(s): 1
AB
     The scarce bioavailability of \beta -interferon (
     IFN-β ) after i.m. administration is probably due
     either to the binding of IFN-\beta to the
     interstitial matrix, or to lymphatic absorption and/or to local breakdown
     by lysosomal proteinases from muscle. In this work, the authors first
     showed that after i.m. injection, the apparent bioavailability of natural
     human IFN-\beta is about 10% of that of
     recombinant IFN-\alpha 2 and then they
     evaluated the effects of proteinase inhibitors and albumin on
     IFN-β incubated at 37° with muscle
     homogenate. IFN biol. activity decreased spontaneously by about 20% after
     incubation for 6 h at 37° in Hanks' solution, but it was almost
     completely lost after incubation with muscle homogenate. Proteinase
     inhibitors (\alpha1-antitrypsin, \alpha2-macroglobulin, aprotinin,
     soybean trypsin inhibitor, leupeptin, EP-459, and EP-475) failed to block
     the inactivation of IFN-\beta by muscle proteinases,
     whereas albumin exerted a partial but consistent protection.
ST
     interferon beta bioavailability muscle albumin
     ; proteinase inhibitor interferon beta bioavailability
ΙT
     Muscle, metabolism
        (interferon-\beta of humans inactivation by,
        albumin and proteinase inhibitors effect on)
ΙT
     Albumins, biological studies
     RL: BIOL (Biological study)
        (muscle inactivation of human interferon-β
        inhibition by)
ΙT
     Interferons
     RL: BIOL (Biological study)
        (β, muscle inactivation of human, albumin and
        proteinase inhibitors effect on)
     138674-34-7, Cysteine proteinase inhibitor 139691-92-2, Serine
ΙT
     proteinase inhibitor
     RL: BIOL (Biological study)
        (muscle inactivation of human interferon-\beta
```

response to)

٠٠٠ المتاريخ

والمتقرقة

```
L66 ANSWER 21 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
     1991:478932 HCAPLUS
AN
DN
     115:78932
ΕĎ
     Entered STN: 23 Aug 1991
     Stable formulations of lipophilic recombinant proteins
ΤI
IN
     Fernandes, Peter M.; Taforo, Terrance
     Cetus Corp., USA
PA
     U.S., 20 pp. Cont.-in-part of U.S. Ser. No. 752,403.
SO
DT
     Patent
LA
     English
     ICM A61K037-02
IC
     ICS A61K045-02
NCL
     424085200
     63-6 (Pharmaceuticals)
     Section cross-reference(s): 16
FAN.CNT 3
    PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
                                                            -----
                                           ______
                           _____
     US 4992271
                            19910212
                                           US 1985-775751
                                                            19850913 <--
                      Α
                           19840731
                                           US 1983-495896
                                                            19830518 <--
     US 4462940
                      Α
     CA 1339707
                      A1 19980310
                                           CA 1986-516417
                                                            19860820 <--
                      A1
                          19870319
                                           AU 1986-62642
                                                            19860912 <--
     AU 8662642
     AU 590896
                      В2
                            19891123
                                         EP 1986-307070
     EP 215658
                      A2
                            19870325
                                                            19860912 <--
     EP 215658
                      А3
                            19890208
     EP 215658
                       В1
                            19940601
         R: AT, BE, CH, DE, FR, GB, IT, LI, NL, SE
     AT 106247
                      E
                            19940615
                                           AT 1986-307070
                                                            19860912 <--
     JP 62067032
                      Α2
                            19870326
                                           JP 1986-215063
                                                            19860913 <--
     JP 06004542
                      В4
                            19940119
     US 5643566
                      Α
                            19970701
                                           US 1995-474769
                                                            19950607 <--
PRAI US 1982-422421
                           19820923
                                     <--
     US 1983-495896
                            19830518
                                     <--
     US 1984-592077
                            19840323
                                     <--
     US 1985-752403
                            19850705. <--
     US 1985-775751
                            19850913
                                     <--
     EP 1986-307070
                            19860912
                                     <--
     US 1986-923425
                            19861027
                                      <--
     US 1992-865411
                            19920507
                                      <--
     US 1994-266832
                            19940628
                                     <--
     An improved process for recovering and purifying lipophilic
AB
     recombinant proteins such as human \beta -
     interferon and interleukin-2 (IL-2) from their hosts yields a
     protein preparation which is formulated into a stable pharmaceutical
composition
     having a therapeutically effective amount of the biol. active
     recombinant lipophilic protein dissolved in a nontoxic, inert,
     therapeutically compatible aqueous based carrier medium at a pH of 6.8 to 7.8.
     The medium also contains a stabilizer for the protein, such as human serum
     albumin and human plasma protein fraction. IL-2 produced by
     recombinant Escherichia coli was purified by a series of steps and
     formulated with human serum albumin (final concentration 2.5%) at pH
ST
     interleukin Escherichia albumin stabilizer; interferon
     recombinant albumin formulation
ΙT
     Escherichia coli
        (beta-interferons and interleukin 2 from)
     Proteins, biological studies
ΙT
     RL: BIOL (Biological study)
        (of blood plasma, as stabilizers for recombinant interleukin
        2-containing pharmaceutical compns.)
```

```
ΙT
     Pharmaceutical dosage forms
        (of recombinant \beta -interferons and
        interleukin 2, stabilizers in, albumins and sugars as)
     Albumins, biological studies
ΤТ
     RL: BIOL (Biological study)
        (stabilizers, for recombinant interleukin 2-containing
        pharmaceutical compns.)
     Lymphokines and Cytokines
ΙT
     RL: BIOL (Biological study)
        (interleukin 2, recombinant, from Escherichia coli,
        stabilized formulations of, albumins and sugars in)
ΙT
     Interferons
     RL: BIOL (Biological study)
        (β , recombinant, from Escherichia coli,
        stabilized formulations of, albumins and sugars in)
ΙT
     69-65-8, Mannitol
     RL: BIOL (Biological study)
        (stabilizer, for recombinant interleukin-2 containing
        pharmaceutical composition)
     50-99-7, Dextrose, biological studies
ΙT
     RL: BIOL (Biological study)
        (stabilizer, for recombinant \beta -
        interferon-containing pharmaceutical composition)
L66 ANSWER 22 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
     1990:153049 HCAPLUS
AN
DN
     112:153049
ΕD
     Entered STN: 28 Apr 1990
TΤ
     Use of human serum albumin signal peptide in recombinant
     protein manufacture and secretion with yeast
     Hayasuke, Naofumi; Nakagawa, Yukimitsu; Ishida, Yutaka; Okabayashi, Ken;
IN
     Murakami, Kohji; Tsutsui, Kiyoshi; Ikegaya, Kazuo; Minamino, Hitoshi;
     Ueda, Sadao; et al.
PA
     Green Cross Corp., Japan
     Eur. Pat. Appl., 35 pp.
SO
     CODEN: EPXXDW
DT
     Patent
     English
LA
     ICM C12N015-00
     ICS C12P021-00
CC
     3-4 (Biochemical Genetics)
FAN.CNT 1
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
     PATENT NO.
     ______
                      ____
                           -----
                                           -----
                                                            -----
                                           EP 1988-107087
                                                            19880503 <--
     EP 319641
                     A1
                            19890614
                            19930922
     EP 319641
                      В1
        R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE
                                           JP 1988-103339
                                                            19880426 <--
     JP 02167095
                      A2
                           19900627
                            19980827
     JP 2791418
                       В2
     CA 1326217
                      Α1
                            19940118
                                           CA 1988-565766
                                                            19880503 <---
                                           ES 1988-107087
                      Т3
                            19941116
                                                            19880503 <--
     ES 2059428
                      В1
                            19970414
                                           KR 1988-5553
                                                            19880513 <--
     KR 9705250
                                                            19950522 <--
     US 5503993
                      Α
                            19960402
                                           US 1995-445783
PRAI JP 1987-306674
                      Α
                            19871202
                                      <--
     JP 1988-45605
                      Α
                           19880226
                                      <--
     US 1988-190553
                       В1
                            19880505
                                      <--
     US 1992-913785
                       В1
                            19920630
                                      <--
     MARPAT 112:153049
OS
     A method for producing and secreting proteins with yeast comprises
     transformation of the yeast with a chimeric gene for a human
    'albumin signal peptide and the coding sequence for the desired
```

protein and expression of the gene. Plasmid pNH008, containing the GAL1

promoter linked to a synthetic human serum albumin signal

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sequence fused to the mature human serum albumin gene and the pho5 terminator, was constructed. Saccharomyces cerevisiae AH22 transformed with this plasmid produced 160 mg ${\tt albumin}/{\tt L}$ culture medium after 48 h incubation. protein secretion yeast albumin signal peptide; Saccharomyces ST human albumin manuf secretion Saccharomyces cerevisiae ΙT (human serum albumin manufacture and secretion with, albumin signal peptide in) Molecular cloning ΙT (in yeast, human serum albumin signal sequence in) Albumins, preparation ΙT RL: PREP (Preparation) (manufacture of, of human, with yeast, human serum albumin signal peptide in) ΙT Lymphokines and Cytokines RL: PROC (Process) (manufacture of, with yeast, human serum albumin signal peptide in) IT Protein sequences (of albumin signal peptide analogs, of human) ΙT Yeast (recombinant protein secretion from, signal peptide of human serum **albumin** in) Deoxyribonucleic acid sequences ΙT (albumin-specifying, signal peptide analog, of human) IT Gene and Genetic element RL: BIOL (Biological study) (chimeric, for signal sequence of human serum albumin and desired protein, expression in yeast of, protein secretion in relation to) Plasmid and Episome ΙT (pNH008, chimeric human serum albumin signal peptide-albumin gene on, expression in Saccharomyces cerevisiae of, albumin secretion in relation to) Peptides, biological studies ΙT RL: BIOL (Biological study) (signal, of human serum albumin, protein secretion from recombinant yeast using) ΙT Gene and Genetic element, animal (signal sequence, of human serum albumin gene, protein secretion from yeast in relation to) Interferons TΤ RL: PROC (Process) $(\alpha$, manufacture of, with yeast, human serum albuminsignal peptide in) IT Interferons RL: PROC (Process) $(\beta$, manufacture of, with yeast, human serum ${\tt albumin}$ signal peptide in) IT Interferons RL: PROC (Process) $(\gamma, \text{ manufacture of, with yeast, human serum albumin signal})$ peptide in) 125677-92-1P 125677-93-2P 125677-94-3P 125677-91-0P IT 125677-90**-**9P 125677-95-4P RL: PREP (Preparation) (human serum albumin signal peptide derivative, recombinant protein manufacture and secretion with yeast in relation to) 125677-89-6P ΙT RL: PREP (Preparation)

(human serum albumin signal peptide, recombinant

```
protein manufacture and secretion with yeast in relation to)
     9001-27-8P, Factor VIII 9002-72-6P, Growth hormone
IT
                                  9039-53-6P, Urokinase
                                                           11096-26-7P,
     Insulin, biological studies
     Ervthropoietin
                     62683-29-8P, Colony-stimulating factor
                                                               85637-73-6P,
    Atriopeptin
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (manufacture and secretion of, with yeast, human serum albumin
        signal peptide in relation to)
    126115-99-9P
ΙT
    RL: PREP (Preparation)
        (nucleotide sequence encoding human serum albumin signal
        peptide, recombinant protein manufacture and secretion with yeast
        in relation to)
    ANSWER 23 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
ΑN
    1989:639534 HCAPLUS
    111:239534
DN
ED
    Entered STN: 23 Dec 1989
    Pharmaceutical compositions containing recombinant
TΙ
     interferon-B
     Taforo, Terrance; Thomson, Jody; Shaked, Ze'ev; Hershenson, Susan;
ΙN
    Thomson, James W.; Stewart, Tracy
PΑ
    Cetus Corp., USA
    PCT Int. Appl., 80 pp.
SO
    CODEN: PIXXD2
DT
    Patent
    English
LA
IC
     ICM A61K047-00
     ICS A61K045-02
     63-6 (Pharmaceuticals)
FAN.CNT 2
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO. DATE
                           -----
                                           _____
PΙ
    WO 8902750
                     A1
                            19890406
                                          WO 1988-US3313
                                                           19880926 <--
        W: AU, DK, JP, NO
        RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE
                            19930202
                                           US 1987-100679
                                                            19870929 <--
    US 5183746
                      Α
    AU 8825351
                            19890418
                                           AU 1988-25351
                                                            19880926 <--
                      Α1
PRAI US 1987-100679
                            19870929
                                     <--
    US 1986-923423
                            19861027
                                     <--
    WO 1988-US3313
                           19880926 <--
    A stable parenteral composition in liquid or lyophilized form comprises a
AB
    recombinant interferon-\beta (IFN-.
    beta.) protein dissolved in an inert carrier medium containing
    nonionic polymeric surfactants as a solubilizer/stabilizer. The
     surfactants include polyoxyethylene sorbitan fatty acid esters, a mixture of
     ethoxylated fatty alc. ethers and lauryl ether, ethoxylated octylphenol, a
    mixture of ethoxylated or propoxylated alcs., polyethylene glycol
    monooleate, ethoxylated phenol, and propylene oxide-ethylene oxide block
     copolymers. The composition further comprises addnl. bulking/stabilizing
     agents, such as dextrose. An IFN-\beta analog
     designated as IFN-\beta ser17 was recovered from
    Escherichia coli culture media and stabilized by adding 0.15% Trycol
    LAL-12 and pH was adjusted to 7.0 with NaOH. A bulking/stabilizing agent,
     i.e., 5% dextrose, was then added and the solution was sterile-filtered,
    aseptically filled into vials, and lyophilized. The IFN-.
    beta. formulations of this invention contain very low levels of
    aggregates and other potentially immunogenic characterisitcs and minimal
    or no strong solubilizing agents, such as SDS, and they are nontoxic and
    have good shelf life.
ST
     interferon beta surfactant solubilizer injection;
     lyophilization interferon beta stability
    Solubilizers
IT
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ور تاليخون

Stabilizing agents

ور والتقالم

55

```
(nonionic surfactants and sugars as, for interferon
        β -containing parenteral compns.)
ΙT
    Albumins, biological studies
    RL: BIOL (Biological study)
        (parenteral interferon-β composition containing
        nonionic surfactants and, as stabilizer)
ΙΤ
    Carbohydrates and Sugars, biological studies
    RL: BIOL (Biological study)
        (parenteral interferon-β composition containing
       nonionic surfactants and, as stabilizers)
ΙT
    Surfactants
        (nonionic, parenteral interferon-\beta composition
        containing, as stabilizers)
IT
     Pharmaceutical dosage forms
        (parenterals, containing \beta -interferons, nonionic
        surfactants and sugars in, as solubilizers/stabilizers)
IT
    Interferons
    RL: BIOL (Biological study)
        (β , parenteral compns. containing, solubilizers/stabilizers
        for, nonionic surfactants and sugars as)
     50-70-4, Sorbitol, biological studies 50-99-7, Dextrose, biological
IT
              56-81-5, Glycerol, biological studies 69-65-8, Mannitol
    87-89-8, Inositol 151-21-3, Sodium dodecyl sulfate, biological studies
    RL: BIOL (Biological study)
        (parenteral interferon-β composition containing
        nonionic surfactants and, as stabilizer)
     9002-92-0, Ethoxylated lauryl alcohol 9002-93-1, Triton X305
IT
     9004-78-8, Ethoxylated phenol 9004-96-0 9005-64-5, Polyoxyethylene
     sorbitan monolaurate 9005-65-6 9036-19-5, Ethoxylated octylphenol
    12616-49-8, Plurafac C17
                               106392-12-5, Propylene oxide-ethylene oxide
    blocker copolymer
    RL: BIOL (Biological study)
        (parenteral interferon-\beta composition containing, as
        stabilizer)
L66 ANSWER 24 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
ΑN
    1989:18548 HCAPLUS
DN
    110:18548
ΕD
    Entered STN: 21 Jan 1989
    Method for treatment of essential (hemorrhagic) thrombocythemia with human 3
TI
    \alpha -interferon
    Delwiche, Francis; Flament-Grivegnee, Jocelyn; Gangji, Diamond; Monsieur,
ΙN
    Rita; Stryckmans, Pierre; Velu, Thierry; Wybran, Joseph
    Boehringer Ingelheim International G.m.b.H., Fed. Rep. Ger.
PΑ
SO
    U.S., 4 pp.
    CODEN: USXXAM
DT
    Patent
LA
    English
    ICM A61K045-02
IC
NCL
    424085000
    1-8 (Pharmacology)
    Section cross-reference(s): 63
FAN.CNT 1
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                            DATE
                     ---- ------ ,
                                        -----
    ______
    US 4743445
                    A 19880510
                                          US 1985-758729
                                                           19850725 <--
PΙ
PRAI US 1985-758729
                          19850725 <--
    Essential thrombocythemia is treated by administration of an effective
    amount of human \alpha -interferon. Patients with
    essential thrombocythemia were given i.m. injections of 5 + 106 IU
     recombinant human interferon-\alpha 2 (Arg)
```

(I)/day for 30 days. After 15 days, the dose was doubled if the results

of the treatment were insufficient. After 30 days, the same dose was given twice a week as a maintenance dose. In all patients the number of thrombocytes returned to normal. A parenteral formulation comprises I 5 + 106 IU, isotonic phosphate buffer (pH 7) q.s., human serum albumin 20.0 mg, and water for injection 1.0 mL. STessential thrombocythemia alpha interferon ΙT Blood platelet (α -interferon of human effect on) ΙT Blood platelet (disease, essential thrombocythemia, treatment of, with α -interferon of human) IT Interferons RL: BIOL (Biological study) $(\alpha$, essential thrombocythemia treatment with, of human) 118104-04-4 RL: BIOL (Biological study) (essential thrombocythemia treatment with) ANSWER 25 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN L66 ΑN **1988:562850** HCAPLUS DN 109:162850 EDEntered STN: 12 Nov 1988 TIRecombinant human interferon alpha-2a: delivery to lymphoid tissue by selected modes of application ΑU Supersaxo, Andreas; Hein, Wayne; Gallati, Harald; Steffen, Hans CS Preclin. Dev., F. Hoffmann-La Roche und Co. Ltd., Basel, Switz. SO Pharmaceutical Research (1988), 5(8), 472-6 CODEN: PHREEB; ISSN: 0724-8741 DT Journal LA English CC 1-2 (Pharmacology) Following s.c. or injection device (i.d.) administration, AB recombinant human interferon α -2a (rIFN α -2a) of mol. weight 19,000 was absorbed mainly by the lymphatics. This results in high rIFN α -2a levels in the lymphoid tissue which drains the application site, while blood plasma levels are relatively low. The maximum measured concns. of rIFN α -2a in the efferent popliteal lymph varied by a factor of 105 between intradermal/s.c. and i.v. administration and was affected neither by the infusion rate nor by the coadministration of albumin. This may help to improve the mode of administration and therapeutic efficacy of protein drugs whose targets are lymphoid cells. interferon α 2a delivery lymph gland ST ΙT Lymphatic system (interferon α -2a absorption by, after parenteral administrations) ΙT Albumins, biological studies RL: BIOL (Biological study) (interferon α -2a delivery to lymphoid tissue in relation to) ΙT Lymph gland (interferon α -2a delivery to, parenteral administration routes for) ΙT Interferons RL: BIOL (Biological study) $(\alpha - 2a, delivery to lymphoid tissue of$ recombinant, parenteral administration routes for) ANSWER 26 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN L66 1987:583557 HCAPLUS ΑN DN 107:183557 Entered STN: 14 Nov 1987 ΕD

Improved formulation for recombinant β -

ور والترب

٠ : الكنوب

ΤI

```
interferon with protein or sugar stabilizer
     Hanisch, Wolfgang Helmut; Taforo, Terrance; Fernandes, Peter Michael
ΙN
PΑ
     Cetus Corp., USA
     Eur. Pat. Appl., 34 pp.
SO
     CODEN: EPXXDW
DT
     Patent
LA
     English
IC
     ICM A61K045-02
     ICS A61K047-00; C07K003-02; C12P021-02
     63-6 (Pharmaceuticals)
CC
     Section cross-reference(s): 3
FAN.CNT 3
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                                            _____
                            _____
                                            EP 1986-307070
                                                             19860912 <--
                      A2
                            19870325
PΙ
     EP 215658
     EP 215658
                            19890208
                       Α3
     EP 215658
                      В1
                           19940601
         R: AT, BE, CH, DE, FR, GB, IT, LI, NL, SE
     US 4992271
                                            US 1985-775751
                                                             19850913 <--
                     Α
                            19910212
     AT 106247
                                                             19860912 <--
                            19940615
                                            AT 1986-307070
                       E
PRAI US 1985-775751
                            19850913
                                      <--
     US 1982-422421
                            19820923
                                      <--
     US 1983-495896
                            19830518
                                      <--
     US 1984-592077
                            19840323
                                      <--
     US 1985-752403
                            19850705
                                      <--
     EP 1986-307070
                            19860912
     Recombinant β-human interferon (.beta
AΒ
     .-HIFN) is dissolved in a non-toxic, inert, therapeutically compatible aqueous
     carrier, at a pH of 2-4. The solution contains a stabilizer for the
     \beta\textsc{-HIFN}, particularly human plasma protein fraction, human serum
     albumin, or mannitol. This formulation results in very low sodium
     dodecyl sulfate levels. \beta -Interferon 0.25 mg/mL
     was formulated using 2.5% plasma protein fraction at pH 3-4, incubated
     15-45 min.; the pH was adjusted to 7.3-7.5. At this pH, the solns. were
     very clear. The use of 5.0% human serum albumin also gave clear
     solns., whereas 2.5% HSA resulted in slightly hazy solns.
     interferon formulation protein solubilization; stabilizer
ST
     recombinant beta interferon
     Albumins, biological studies
ΤТ
     RL: BIOL (Biological study)
        (human, stabilizer for recombinant \beta-human
        interferon)
     Proteins, specific or class, biological studies
TΨ
     RL: BIOL (Biological study)
        (of blood plasma, as stabilizer for recombinant \beta-human
        interferon)
IT
     Recombination, genetic
        (of \beta -interferon, purification and formulation for)
TΤ
     Interferons
        (\beta -, recombinant, stabilization of, in
        formulation)
     151-21-3, Sodium dodecyl sulfate, biological studies
IT
     RL: PRP (Properties)
        (reduced levels of, in formulations of \beta -
        interferon)
ΙT
     50-99-7, Dextrose, biological studies
                                              69-65-8, Mannitol
     RL: BIOL (Biological study)
        (stabilizer, for recombinant \beta -
        interferon-containing pharmaceutical composition)
    ANSWER 27 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
     1987:464710 HCAPLUS
ΑN
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107:64710

DN

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Entered STN: 21 Aug 1987
ED
ΤI
     Potency stability of recombinant (serine-17) human
     interferon-β
     Geigert, John; Ziegler, Diana L.; Panschar, Barbara M.; Creasey, Abla A.;
AU
     Vitt, Charles R.
CS
     Dep. Tech. Dev., Cetus Corp., Emeryville, CA, 94608, USA
     Journal of Interferon Research (1987), 7(2), 203-11
SO
     CODEN: JIREDJ; ISSN: 0197-8357
DT
     Journal
     English
LA
CC
     63-3 (Pharmaceuticals)
     The antiviral activity of Escherichia coli-derived (serine-17) human
AΒ
     interferon-\beta , formulated with human serum
     albumin, is stable for 2 yr when lyophilized and stored under
     refrigeration. This product shows an Arrhenius line fit for the stability
     of its activity when tested at multiple isothermal temps. (25-80°).
     In both isothermal and non-isothermal elevated temperature studies, increasing
     the level of human serum albumin in the formulation results in
     increased thermal stability.
     interferon serine 17 recombinant formulation stability
ST
     Kinetics of decomposition
        (of recombinant human β -interferon
        in albumin formulation)
     Albumins, uses and miscellaneous
ΙT
     RL: USES (Uses)
        (β -interferon recombinant serine-17
        stabilization by formulation with human)
ΙT
     Interferons
        (\beta -, stability of recombinant serine-17, in
        human serum albumin formulation)
    ANSWER 28 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN
L66
     1986:174635 HCAPLUS
ΑN
DN
     104:174635
ED
     Entered STN: 17 May 1986
     Interferon solubilization with amino acids
TI
     Kato, Yasuki; Hayakawa, Eiji; Furuya, Kunitoshi; Kondo, Akira
ΙN
     Kyowa Hakko Kogyo Co., Ltd., Japan
PΑ
SO
     Eur. Pat. Appl., 14 pp.
     CODEN: EPXXDW
DT
     Patent
     English
LA
     ICM A61K045-02
IC
     63-3 (Pharmaceuticals)
     Section cross-reference(s): 15
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                            DATE
                      ----
                                           -----
                                                           _____
     EP 163111
                            19851204
                                           EP 1985-104849
                                                            19850422 <--
                      A2
     EP 163111
                     А3
                            19870930
                            19901003
     EP 163111
                     В1
        R: DE, FR, GB, IT
     JP 60243028
                     A2
                            19851203
                                           JP 1984-86972
                                                            19840428 <--
     JP 05058000
                       B4
                            19930825
     CA 1264665
                       Α1
                            19900123
                                           CA 1985-479841
                                                            19850423 <--
     US 4675183
                      Α
                            19870623
                                           US 1985-726971
                                                            19850425 <--
PRAI JP 1984-86972
                            19840428
                                     <--
     Interferon is solubilized by addition of 5 + 10-6 - 5 +
     10-3 mol amino acid/106 units interferon. The amino acid may be
     arginine, histidine, lysine, hydroxylysine, ornithine, glutamine,
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 γ -aminobutyric acid, ϵ -aminocaproic acid, or a salt of these

compds. Thus, 5 mg serum albumin, 5 mg NaCl, 30 mg arginine-HCl, and 3 + 106 units of γ - interferon were

المنتقع المنتقع

ST

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SO

DT

LACC

ST

IT

(albumin effect on recombinant $\alpha 2$ -

mixed with 2 mL H2O, and freeze-dried. The product was dissolved in 5 mL H2O, held 6 h at 25°, and the absorbance was measured at 400 nm. The amount of γ - interferon that remained in solution was 98%. This solubilization may be used to facilitate the isolation and purification of interferon produced by recombinant DNA technol. interferon solubilizer amino acid; arginine interferon solubilization Solubilizers (amino acids, for interferon) Amino acids, uses and miscellaneous RL: PRP (Properties) (interferons solubilization by) Interferons $(\alpha$ -, solubilization of, with amino acids) Interferons $(\beta$ -, solubilization of, with amino acids) Interferons $(\gamma$ -, solubilization of, with amino acids) 56-87-1, properties 60-32-2 70-26-8 56-85-9, properties properties 74-79-3, properties 657-27-2 1119-34-2 1190-94-9 60259-81-6 2835-81-6 RL: PRP (Properties) (interferons solubilization by) ANSWER 29 OF 29 HCAPLUS COPYRIGHT 2004 ACS on STN L66 1986:86802 HCAPLUS 104:86802 Entered STN: 22 Mar 1986 The lymphatic route - II. Pharmacokinetics of human recombinant interferon- α 2 injected with albumin as a retarder in rabbits Bocci, Velio; Muscettola, Michela; Naldini, Antonella; Bianchi, Enrica; Segre, Giorgio Inst. Gen. Physiol., Univ. Siena, Siena, 53100, Italy General Pharmacology (1986), 17(1), 93-6 CODEN: GEPHDP; ISSN: 0306-3623 Journal English 15-5 (Immunochemistry) An investigation was conducted to define whether multisite s.c. administration in unanesthetized, unrestrained rabbits of human recombinant interferon- α 2 (rec. IFN- α 2) either in saline, human albumin (ALB) solution (4, 7, and 10% final concns.), or in a solution containing 75 units of hyaluronidase, modified the pharmacokinetic parameters calculated from the IFN plasma level. Plasma disappearance rates of rec. IFN-. alpha.2 were measured in rabbits after i.v. administration and the kinetics was adequately represented by a 3-compartment mammillary model. This model was the basis for evaluating the absorption and distribution of rec. IFN- α 2 after s.c. administration. The increase of ALB concentration (from 4 to 10%) caused a significant reduction plasma IFN maximum clearance, while both the mean residence time and the release time of IFN increased linearly with the ALB concentration The data support the postulation that s.c. administration of albumin acts as an interstitial fluid expander and may favor absorption of IFN via lymphatics rather than blood capillaries. Improvement of therapeutic index of IFN by using this route remains to be shown in clin. trials. interferon alpha pharmacokinetics albumin Lymphatic system

interferon pharmacokinetics in relation to, of humans and laboratory

animals)

IT Blood plasma

(α 2- interferon pharmacokinetics in, albumin effect on, in humans and laboratory animals)

IT Albumins

RL: BIOL (Biological study)

(α 2- interferon pharmacokinetics response to, of humans and laboratory animals)

IT Interferons

RL: BIOL (Biological study)

(α 2-, pharmacokinetics of recombinant

, albumin effect on, of humans and laboratory animals)

=> => fil wpix

FILE 'WPIX' ENTERED AT 16:25:05 ON 02 FEB 2004

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FILE LAST UPDATED:

· 28 JAN 2004

<20040128/UP>

`<<<

MOST RECENT DERWENT UPDATE: 200407

00407 <200407/DW>

DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

- >>> NEW WEEKLY SDI FREQUENCY AVAILABLE --> see NEWS <
- >>> SLART (Simultaneous Left and Right Truncation) is now
 available in the /ABEX field. An additional search field
 /BIX is also provided which comprises both /BI and /ABEX <<</pre>
- >>> PATENT IMAGES AVAILABLE FOR PRINT AND DISPLAY <<<
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- >>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://thomsonderwent.com/coverage/latestupdates/ <<<
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>>> ADDITIONAL POLYMER INDEXING CODES WILL BE IMPLEMENTED FROM DERWENT UPDATE 200403.

THE TIME RANGE CODE WILL ALSO CHANGE FROM 018 TO 2004.

SDIS USING THE TIME RANGE CODE WILL NEED TO BE UPDATED.

FOR FURTHER DETAILS: http://thomsonderwent.com/chem/polymers/ <<<

=> d all abeq tech abex tot

L88 ANSWER 1 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2003-421048 [39] WPIX

DNC **C2003-110745**

TI New hybrid polypeptide, useful for sequestering and/or purifying a polypeptide of interest.

DC B04 D16

IN THOMAS, T; TILLETT, D

PA (PROT-N) PROTIGENE PTY LTD

CYC 101

- -

PI WO 2003018616 A1 20030306 (200339) * EN 66p C07K001-14

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR IE IT KE LS LU

MC MW MZ NL OA PT SD SE SK SL SZ TR TZ UG ZM ZW

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ADT WO 2003018616 A1 WO 2002-AU1159 20020827

PRAI AU 2001-7298 20010827

ICM C07K001-14

C07K001-36; C07K019-00; C12N009-00; C12N015-63

AB WO2003018616 A UPAB: 20030619

> NOVELTY - A hybrid polypeptide comprises a polypeptide of interest linked to a polymerizable polypeptide, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- sequestering and/or purifying a polypeptide of interest;
- (2) a hybrid nucleic acid comprising a nucleic acid encoding the hybrid polypeptide;
- (3) a library comprising several hybrid nucleic acids, polypeptides or vectors;
 - (4) a vector comprising the hybrid nucleic acid;
- (5) a cell transformed or transfected with the hybrid nucleic acid or vector; and
 - (6) purifying a polypeptide of interest.

UPTX: 20030619

USE - The hybrid polypeptide is useful for sequestering and/or purifying a polypeptide of interest (claimed). Dwq.0/9

CPI FS

AB; DCN FΑ MC CPI: B04-B04C; B04-C01; B04-E08; B04-F0100E; B04-G01; B04-H01; B04-H02B; B04-H04; B04-H05; B04-H19; B04-J01; B04-J02; B04-J05; B04-J10; B04-L04; B04-L05; B04-L06; B04-L07; B04-N03; B04-N04; B04-N06; B04-N08; B11-B; D05-C11; D05-H12A; D05-H12E; D05-H13; D05-H14; D05-H17C

TECH

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TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Polypeptide: The hybrid polypeptide is produced in vivo. It is linked to a support, comprising the polymerizable polypeptide. The support polymerizable polypeptide comprises a polymerizable polypeptide identical to the hybrid polypeptide, or its variant. The polypeptide of interest is linked to the polymerizable polypeptide by fusing the polypeptide of interest directly to the polymerizable polypeptide or by a linker polypeptide. It is prokaryotic or eukaryotic in origin. It is a synthetic polypeptide. It comprises endonuclease, a methylase, an oxidoreductase, a transferase, a hydrolase, a lysase, an isomerase, a ligase, a storage polypeptide, a fertitin, an ovalbumin, a transport protein, hemoglobin, serum albumin or ceruloplasmin, an antigen, an antigenic determinant for use in the preparation of vaccines or diagnostic agents, a protective protein, a defense protein, thrombin, fibrinogen, binding proteins, antibodies, immunoglobulins, a human growth hormone, somatostatin, prolactin, estrange, progesterone, melanocyte, thyrotropin, calcitonin, gonadotropin, insulin, a hormone identified as being involved in the immune system, interleukin 1, interleukin 2, colony simulating factor, macrophage-activating factor, interferon, a structur al element, collagen, elastin, alpha-keratin, glyco-protein, virus-protein and muca-protein. The linker polypeptide comprises a recognition site for a proteolytic agent and a multiple cloning site. It also comprises a spacer polypeptide of sufficient length to allow or enhance cleavage of the polypeptide of interest from the polymerizable polypeptide, or to avoid unfavorable steric interference between the polypeptide of interest and the polymerizable polypeptide.

The recognition site comprises an amino acid sequence consisting of:

- (a) Leu-Glu-VaI-Leu-Phe-Gln-Gly-Pro;
- (b) Leu-Val-Pro-Arg-Gly-Ser;

- (c) Ile-Glu-Gly-Arg; or
- (d) Asp-Asp-Asp-Asp-Lys.

The chemical capable of proteolytic activity is cyanogen bromide. The polypeptides are linked by antibody interaction, which is achieved by:

- (a) attaching an antibody specific for the polypeptide of interest to the polymerizable polypeptide; or
- (b) using a bi-specific antibody directed to both the polypeptide of interest and the polymerizable polypeptide.

The polymerizable polypeptide is a polypeptide that naturally polymerizes with itself. It is tubulin or actin. It is an FtsZ or Escherichia coli FtsZ protein or its variant. The variant Escherichia coli FtsZ protein comprises replacement of the aspartate residue at position 212 of the protein with a cysteine or asparagine residue. The variant FtsZ protein comprises a mutation selected from replacement of alanine by threonine at position 70, replacement of aspartate by alanine at position 209 or replacement of aspartate by alanine at position 269. The polymerizable polypeptide requires an intermediary polypeptide or other molecule in order to polymerize.

Preferred Method: Sequestering and/or purifying a polypeptide of interest comprises polymerizing the hybrid polypeptide under controlled chemical and/or physical conditions. It is polymerized by a change in temperature and by the addition of an agent that induces polymerization. The polymerization inducing agent is GTP, ATP and/or a cation. The cation comprises magnesium, calcium, nickel, cobalt, zinc or manganese. The polymerized hybrid polypeptide is purified by a first purification step, which may be the only purification step or may be followed by further purification steps. The first purification step purifies the polymerized hybrid polypeptide by physical techniques discriminating on the basis of size and/or weight. The polymerized hybrid polypeptide is also purified by centrifugation, differential sedimentation, filtration, dialysis and/or flow sorting, where the polymerized hybrid polypeptide is isolated. After the first purification step the polymerized hybrid polypeptide is dissociated. The dissociation is achieved by removal of the agent which induces polymerization and/or incubation of the polymerized hybrid polypeptide at a suitable temperature. The dissociated hybrid polypeptide is purified by a second purification step, which comprises purification of the hybrid polypeptide on the basis of size and/or weight. The polymerization, dissociation and purification of the polymerizable hybrid polypeptide are repeated so that substances larger and smaller than the hybrid polypeptide are removed. The polymerizable polypeptide is cleaved from the polypeptide of interest by a proteolytic agent, which does not substantially interfere with the biological or chemical activity of the polypeptide of interest or the polymerizable polypeptide. After the cleavage of the polypeptide of interest from the polymerizable polypeptide, the protease hybrid polypeptide is polymerized. The proteolytic agent comprises 3C-protease from a human rhinovirus type 14 (HRV protease 3C), thrombin, Factor Xa, enterokinase and a chemical capable of proteolytic activity. It is linked to a polymerizable polypeptide to form a protease hybrid polypeptide. The polymerizable polypeptide to which the protease is linked is identical to the polymerizable polypeptide to which the polypeptide of interest is linked, or is a variant of it.

Purifying a polypeptide of interest comprises:

- (a) expressing the hybrid nucleic acid in a cell to produce a hybrid polypeptide comprising the polypeptide of interest and a polymerizable polypeptide;
- (b) polymerizing the hybrid polypeptide;
- (c) purifying the polymerized hybrid polypeptide;
- (d) cleaving the polypeptide of interest from the polymerizable polypeptide; and
- (e) purifying the polypeptide of interest.

ABEX

مروانيتن

UPTX: 20030619

EXAMPLE - No suitable example given.

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L88 ANSWER 2 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN
AN
     2002-179329 [23]
                        WPIX
CR
     2001-602931 [68]
DNC
     C2002-055553
ΤI
     New albumin fusion proteins with extended shelf life, useful for
     treating leukemia, warts, hepatitis, multiple sclerosis and AIDS,
     comprises therapeutic protein fused to albumin.
     B04 D16
DC
ΙN
     BALLANCE, D J; PRIOR, C P; SADEGHI, H; SLEEP, D; TURNER, A J
PΑ
     (DELZ) DELTA BIOTECHNOLOGY LTD; (PRIN-N) PRINCIPIA PHARM CORP; (BALL-I)
     BALLANCE D J; (PRIO-I) PRIOR C P; (SADE-I) SADEGHI H; (SLEE-I) SLEEP D;
     (TURN-I) TURNER A J
CYC
     96
     WO 2001079271 A1 20011025 (200223) * EN 294p
PΙ
                                                     C07K014-00
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            LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD
            SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
     AU 2001061024 A 20011030 (200225)
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     EP 1278767
                   A1 20030129 (200310)
                                         ΕN
                                                     C07K014-00
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            RO SE SI TR
     US 2003199043 A1 20031023 (200370)
                                                     C12P021-02
                                             453p
     JP 2003530839 W 20031021 (200373)
                                                     C12N015-09
ADT WO 2001079271 A1 WO 2001-US12009 20010412; AU 2001061024 A AU 2001-61024
     20010412; EP 1278767 A1 EP 2001-934875 20010412, WO 2001-US12009 20010412;
     US 2003199043 A1 Provisional US 2000-229358P 20000412, Provisional US
     2000-199384P 20000425, Provisional US 2000-256931P 20001221, US
     2001-832501 20010412; JP 2003530839 W JP 2001-576866 20010412, WO
     2001-US12009 20010412
FDT AU 2001061024 A Based on WO 2001079271; EP 1278767 A1 Based on WO
     2001079271; JP 2003530839 W Based on WO 2001079271
PRAI US 2000-256931P 20001221; US 2000-229358P 20000412; US 2000-199384P
     20000425; US 2001-832501
                                20010412
TC:
     ICM C07K014-00; C12N015-09; C12P021-02
          A61K038-00; A61K038-16; A61K038-21; A61K038-43; A61K038-46;
          A61K038-48; A61K038-55; A61K039-395; A61K047-48; A61P001-16;
          A61P015-00; A61P017-12; A61P025-28; A61P031-12; A61P031-14;
          A61P031-18; A61P031-20; A61P035-00; A61P035-02; C07H021-04;
          CO7KO14-52; CO7KO14-56; CO7KO14-745; CO7KO14-75;
          C07K014-76; C07K014-765; C07K014-81; C07K016-00;
          C07K019-00; C12N001-19; C12N001-21; C12N005-06; C12N005-10;
          C12N009-14; C12N009-74; C12N009-99; C12N015-00
     WO 200179271 A UPAB: 20031112
AΒ
     NOVELTY - An albumin fusion protein (I) comprising:
          (a) a therapeutic protein (X) and albumin (A) containing a
     fully defined sequence (S1) of 585 amino acids as given in the
     specification;
          (b) X and a fragment or variants of S1, where the fragment or
     variants has albumin activity; or
          (c) a fragment or variant of X and A, where the fragment or variant
     has a biological activity of X, is new.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
     following:
          (1) an albumin fusion protein (II) comprising a peptide
     inserted into A comprising amino acids 54-61, 76-89, 92-100, 170-176,
     247-252, 266-277, 280-288, 362-368, 439-447, 462-475, 478-486 or 560-566
     of S1;
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(2) an **albumin** fusion protein (III) comprising a single chain antibody or its portion and A or its fragment or variant;

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- (3) a composition comprising any of (I)-(III) and a pharmaceutically active carrier;
 - (4) a kit comprising the composition;
- (5) treating a disease or disorder that is modulated by X in a patient comprising administering any of (I)-(III);
- (6) extending the shelf life of X comprising fusing X or its fragment or variant to A or its fragment or variant, sufficient to extend the shelf-life of X compared to the shelf life of X in an unfused state;
- (7) a nucleic acid molecule (IV) comprising a polynucleotide sequence encoding any of (I)-(III);
 - (8) a vector comprising (IV); and
 - (9) a host cell comprising (IV).

ACTIVITY - Cytostatic; dermatological; virucide; anti-HIV; neuroprotective; hepatotropic; antiinflammatory. Tests are described but no results are given in the source material.

MECHANISM OF ACTION - Gene therapy.

USE - The fusion protein is useful for the treatment of hairy cell leukemia, Kaposi's sarcoma, genital warts, anal warts, chronic hepatitis B, chronic non-A, non-B hepatitis, hepatitis C/D, chronic myelogenous leukemia, renal cell carcinoma, bladder carcinoma, ovarian carcinoma, cervical carcinoma, skin cancer, recurrent respirator papillomatosis, non-Hodgkin's lymphoma, cutaneous T-cell lymphoma, melanoma, multiple myeloma, acquired immunodeficiency syndrome (AIDS), multiple sclerosis and glioblastoma. The fusion of albumin extends the shelf life and the in vivo and in vitro biological activity of the therapeutic protein (all claimed).

ADVANTAGE - Therapeutic proteins can be stabilized to extend shelf life and/or retain the protein's activity for extended periods of time in solution, in vivo or in vitro by genetically or chemically fusing the protein to albumin or its fragment or variant. In addition the use of albumin fusion proteins reduces the need to formulate protein solutions with large excesses of carrier proteins to prevent loss of therapeutic protein due to factors such as binding to the container. The extension of shelf life was tested by measuring biological activity (Nb2 cell proliferation) of human albumin-human growth hormone (HA-hGH) fusion protein remaining after incubation in cell culture media for up to 3 weeks at 37 deg. C. At week 3 there was still approx. 95% cell proliferation compared to no activity of unfused hGH (no observed activity by week 2).

Dwg.0/18

FS CPI

TECH

و تأثير

٠٠ تائيتني

FA AB; DCN MC CPI: BO

CPI: **B04-C01G**; B04-E02H; B04-E08; B04-F0100E; B04-G01;

B04-H05A; B04-H19; B04-L05A; B04-N02A; B04-N08;

B14-A02A; B14-A02B1; B14-G01B; B14-H01; B14-N12; B14-N17; B14-S01;

B14-S03A; D05-C12; D05-H12C; D05-H12E; D05-H14; D05-H17C

UPTX: 20020411

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preparation: The fusion proteins can be prepared by standard recombinant techniques. Preferred Fusion Protein: Albumin activity is the ability to prolong the shelf life of X compared to the shelf life of X in an unfused state. Preferably the fragment or variant of (I) comprises amino acids 1-387 of S1. X is chosen from serum cholinesterase, alpha-1 antitrypsin, aprotinin, coagulated complex, von Willebrand factor, fibrinogen, factor VII, factor VIIA activated factor, factor VIII, factor IX, factor X, factor XIII, cl inactivator, antithrombin III, thrombin, prothrombin, apo-lipoprotein, c-reactive protein, protein C, immunoglobulin and preferably interferon (IFN)-alpha. X or its fragment or variant is fused to the N or C-terminus of A. (I)-(III) comprises a first and second X, where the first X is different from the second X. X is separated from A by a linker. The fusion protein has the formula R1-L-R2, R2-L-R1 or R1-L-R2-L-R1, where:

R1 = X

L = peptide linker; and

R2 = A or its fragment or variant.

The in vitro or in vivo activity of X fused to A is greater than the in vitro or in vivo biological activity of X in an unfused state. The protein is expressed in a glycosylation and protease deficient yeast. Alternatively it is expressed by a mammalian cell in culture. The fusion protein further comprises a secretion leader sequence.

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation: The fusion proteins can be produced by standard chemical synthetic techniques.

UPTX: 20020411

ABEX

ADMINISTRATION - 1 microgram/kg/day to 10 mg/kg/day, preferably 0.01-1 mg/kg/day of **albumin** fusion proteins are administered by standard routes.

EXAMPLE - A human albumin-human growth hormone (HA-hGH) fusion protein was prepared. The hGH cDNA was obtained from a human pituitary gland cDNA library by polymerase chain reaction (PCR) amplification. The PCR product was purified and then digested with EcoRl and HindIII. After further purification of the EcoR1-HindIII fragment by gel electrophoresis, the product was cloned into pUC19 digested with EcoR1 and HindIII to give pHGH1. The polylinker sequence of the phagemid pBluescribe (+) (Stratagene) was replaced by inserting an oligonucleotide linker formed by annealing 2 75-mer oligonucleotides between the EcoRl and HindIII sites to form pBST(+). The new polylinker included a unique NotI site. the NotI HA expression cassette of pAYE309 comprising the PRBI promoter, DNA encoding the HA/MFalpha-1 hybrid leader sequence, DNA encoding HA and the ADH1 terminator, was transferred to pBST(+) to form pHA1. The HA sequence was removed from this plasmid by digestion with HindIII followed by religation to form pHA2. Cloning of the hGH cDNA provided the hGH coding region lacking the pro-hGH sequence and the first 8 base pairs (bp) of the mature hGH sequence. In order to construct an expression plasmid for secretion of hGH from yeast, a yeast promoter, signal peptide and the first bp of the hGH sequence were attached to the 5' end of the cloned hGH sequence. The HindIII-SfaNI fragment from pHA1 was attached to the 5' end of the EcoR1/HindIII fragment from pHGHI via 2 synthetic oligonucleotides to generate a double stranded fragment of DNA with sticky ends that can anneal with SfaNI and EcoR1 sticky ends. The HindIII fragment formed was cloned into HindIII digested pHA2 to make pHGH2 such that the hGH cDNA was positioned downstream of the PRBI promoter and HA/MFalpha-1 fusion leader sequence. The NotI expression cassette contained in pHGH2 was cloned into the NotI-digested pSAC35 to make pHGH12. This plasmid comprised the entire 2 micro m plasmid to provide replication functions and the LEU2 gene for selection of transformants. pHGH12 was introduced into S. cerevisiae D88 by transformation and individual transformants were grown for 3 days at 30 degrees C in 10 mL YEPD (1% w/v yeast extract, 2% w/v peptone, 2% w/v dextrose). After centrifugation of the cells, the supernatants were examined by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and were found to contain protein which was of the expected size and recognized by anti-hGHG antiserum on Western blots.

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L88 ANSWER 3 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN
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م ونيتند

AN **2001-616754** [71] WPIX

CR 2001-602931 [68]; 2001-611723 [70]; 2001-616755 [71]; 2001-616756 [71]; 2002-010886 [01]; 2003-810996 [76]; 2004-033644 [03]

DNC **C2001-184720**

Albumin fusion proteins comprising a therapeutic protein and albumin, useful in the treating immune system disorders (e.g. transplant rejection), blood related disorders (e.g. myocardial infarction) and hyperproliferative disorders.

DC B04 D16

IN HASELTINE, W A; ROSEN, C A
PA (HUMA-N) HUMAN GENOME SCI INC

CYC 96

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C12N000-00
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                                                       C12N015-09
                                               469p
     WO 2001079443 A2 WO 2001-US11924 20010412; AU 2001059063 A AU 2001-59063
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      \texttt{JP} \ \ 2003530846 \ \ \texttt{W} \ \ \texttt{JP} \ \ 2001-577427 \ \ \ 20010412, \ \ \texttt{WO} \ \ \ 2001-US11924 \ \ \ \ 20010412 
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     2001079443; JP 2003530846 W Based on WO 2001079443
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          A61K039-395; A61K047-48; A61K048-00; A61P001-16; A61P013-00;
          A61P025-00; A61P031-14; A61P031-18; A61P031-20; A61P035-00;
          A61P035-02; C07K014-47; C07K014-76; C07K019-00;
          C12N001-19; C12N005-10
     WO 200179443 A UPAB: 20040112
AB
     NOVELTY - Albumin fusion proteins (P1) comprising a therapeutic
     protein (T1) (or its fragment or variant having the activity of T1) and
     albumin comprising the 585 amino acid sequence (I) defined in the
     specification (or its fragment or variant having albumin
     activity), are new.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
     following:
          (1) a kit comprising a composition containing P1;
          (2) a method of treating a disease or disorder, preferably modulated
     by T1, in a patient, comprising administering P1;
          (3) a method of extending the shelf-life of T1, comprising fusing T1
     or its fragment or variant, to albumin or its fragment or
     variant, where the shelf-life of T1 or its fragment or variant as part of
     a fused protein is extended when compared to T1 or its fragment or variant
     in an unfused state;
          (4) a nucleic acid (N1) comprising a nucleotide sequence encoding P1;
          (5) a vector comprising N1; and
          (6) a host cell comprising N1.
          ACTIVITY - Cytostatic; antiinflammatory; antileukemic; antiarthritic;
     antirheumatic; immunosuppressive; cardiant; nootropic; neuroprotective;
     antimicrobial; vulnerary.
          To test whether sympathetic neuronal cell viability is supported by
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To test whether sympathetic neuronal cell viability is supported by an albumin fusion protein, the chicken embryo neuronal survival assay (Senaldi, et al., Proc. Natl. Acad., Sci., U.S.A, 96:11458-63 (1998)). Briefly, motor and sympathetic neurons were isolated from chicken embryos, resuspended in L15 medium (with 10% foetal calf serum (FCS), glucose, sodium selenite, progesterone, conalbumin, putrescine and insulin) and Dulbecco's modified Eagles medium (with 10% FCS, glutamine, penicillin, and 25 mM Hepes buffer (pH 7.2)), respectively and incubated at 37 degrees Centigrade in 5% carbon-dioxide in the presence of different concentrations of the purified fusion protein, as well as negative control lacking any cytokine, After 3 days, neuronal survival was determined by evaluation of cellular morphology, and through the use of the colorimetric assay of Mosmann (Mosmann, T., J. Immunol., Methods, 65:55-63 (1983)). Enhanced neuronal cell viability as compared to the controls lacking cytokine is indicative of the ability of the albumin fusion protein to enhance the survival of neuronal cells.

MECHANISM OF ACTION - Gene therapy.

USE - The albumin fusion proteins are also useful in the treatment, prevention, diagnosis, and/or detection of diseases, disorders such as immune system disorders (e.g. transplant rejection), blood related disorders (e.g. myocardial infarction), hyperproliferative disorders (e.g. childhood acute myeloid leukemia), renal disorders (e.g. glomerulonephritis), cardiovascular disorders (e.g. arrhythmias), respiratory disorders (e.g. non-allergic rhinitis), neurological diseases (e.g. Alzheimer's disease), endocrine disorders (e.g. pheocytochroma), reproductive system disorders (e.g. syphilis), infectious diseases (e.g. measles), gastrointestinal disorders (e.g. irritable bowel syndrome) and wound healing.

Dwg.0/15

FS CPI

FA AB; DCN

MC CPI: **B04-C01**; B04-E02F; B04-E08; B04-F0100E; B04-F0200E; B04-F0900E; B04-F1100E; **B04-N02A0E**; B14-A01; B14-A02;

B14-D01; B14-E10; B14-F01; B14-F02; B14-G01; B14-G02; B14-G03;

B14-H01; B14-J01; B14-K01; B14-N10; B14-N17B; B14-S03;

D05-H12B2; D05-H12E; D05-H14A2; D05-H14B2

UPTX: 20011203

TECH

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Fusion Protein: The albumin activity is the ability to prolong the shelf-life of T1 compared to the shelf-life of T1 in an unfused state. The albumin fragment or variant comprises amino acids 1-387 of (I). T1 or its fragment or variant is fused to the C-terminal of the albumin or the C-terminus of the fragment or variant of albumin. Alternatively, T1 or its fragment or variant is fused to the N-terminal of the albumin or the N-terminus of the fragment or variant of albumin. Alternatively, T1 or its fragment or variant is fused to the N-terminus and C-terminus of the albumin, or the N-terminus and C-terminus of the fragment or variant of albumin.

P1 comprises a first T1 or its fragment or variant, and a second T1 or its fragment or variant, where the first T1 is different from the second T1. T1 or its fragment or variant is separated from the **albumin** or the fragment or variant of **albumin** by a linker. Preferably, P1 is of the formula (S1), (S2) or (S3).

R1-L-R2 (S1);

R2-L-R1 (S2); or

R1-L-R2-L-R1 (S3).

Where

R1 = is T1 or its fragment or variant;

L = is a peptide linker; and

R2 = is albumin comprising the sequence of (I), or its fragment or variant.

The shelf-life of the **albumin** fusion protein is greater than the shelf-life of T1 or its fragment or variant in an unfused state. The in vitro or in vivo biological activity of T1 or its fragment or variant, fused to **albumin** or its fragment or variant, is greater than the in vitro or in vivo, respectively, biological activity of T1 or its fragment or variant, in an unfused state.

Alternatively, Pl comprises Tl or its fragment or variant, inserted into an **albumin** comprising the sequence of (I) or its fragment or variant. Preferably, the **albumin** comprises residues 54-61, 76-89, 92-100, 170-176, 247-252, 266-277, 280-288, 362-368, 439-447, 462-475, 478-486, or 560-566 of (I). The portion of **albumin** is sufficient to prolong the shelf-life of Tl, or its fragment or variant, as compared to the shelf-life of Tl, or its fragment or variant in an unfused state.

The portion of **albumin** is sufficient to prolong the in vitro and in vivo biological activity of T1 or its fragment or variant, as compared to the in vitro and in vivo biological activity of T1 or its fragment or

ABEX

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variant, in an unfused state. P1 is non-glycosylated and is expressed in yeast which is glycosylation deficient. The yeast may also be protease deficient. Alternatively, Pl is expressed by a mammalian cell in culture. P1 further comprises a secretion leader sequence. UPTX: 20011203 ADMINISTRATION - The albumin fusion proteins can be administered orally, rectally, parenterally, intracisternally, intravaginally, intraperitoneally, topically, bucally, or as an oral or nasal spray. The dosage is 1 microgram/kg/day to 10 mg/kg/day, preferably 0.01 to 1, mg/kd/day. If given continuously, the albumin fusion protein is typically administered at a dose rate of 1-50 micrograms/kg/hour, either by 1-4 injections per day or by continuous subcutaneous infusions. ANSWER 4 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN **2001-611723** [70] WPIX 2001-602931 [68]; 2001-616754 [71]; 2001-616755 [71]; 2001-616756 [71]; 2002-010886 [01]; 2003-810996 [76]; 2004-033644 [03] C2001-182838 New albumin fusion proteins, useful for treating diseases and disorders such as cancer, comprise therapeutic protein fused to albumin. B04 D16 HASELTINE, W A; ROSEN, C A (HUMA-N) HUMAN GENOME SCI INC 96 WO 2001079442 A2 20011025 (200170)* EN 362p C12N000-00 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR.TT TZ UA UG US UZ VN YU ZA ZW AU 2001064563 A 20011030 (200219) C12N000-00 EP 1276849 A2 20030122 (200315) C12N001-18 ΕN R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR 540p C12N015-09 JP 2003531590 W 20031028 (200373) WO 2001079442 A2 WO 2001-US11850 20010412; AU 2001064563 A AU 2001-64563 20010412; EP 1276849 A2 EP 2001-938994 20010412, WO 2001-US11850 20010412; JP 2003531590 W JP 2001-577426 20010412, WO 2001-US11850 20010412 AU 2001064563 A Based on WO 2001079442; EP 1276849 A2 Based on WO 2001079442; JP 2003531590 W Based on WO 2001079442 PRAI US 2000-256931P 20001221; US 2000-229358P 20000412; US 2000-199384P 20000425 C12N000-00; C12N001-18; C12N015-09 ICM A61K038-00; A61K038-21; A61K039-395; A61K048-00; A61P001-04; A61P001-16; A61P001-18; A61P003-10; A61P005-14; A61P005-40; A61P007-04; A61P007-06; A61P009-00; A61P009-06; A61P009-10; A61P009-12; A61P011-00; A61P011-06; A61P013-00; A61P013-02; A61P013-08; A61P013-12; A61P015-00; A61P015-10; A61P015-18; A61P017-00; A61P017-02; A61P019-00; A61P019-02; A61P019-08; A61P021-00; A61P021-04; A61P025-00; A61P025-08; A61P025-16; A61P025-28; A61P027-02; A61P029-00; A61P031-00; A61P031-12; A61P031-16; A61P031-18; A61P031-22; A61P033-02; A61P033-06; A61P033-12; A61P035-00; A61P035-02; A61P037-00; A61P037-08; A61P039-02; A61P041-00; A61P043-00; C07K014-47; C07K014-76; C07K019-00; C12N001-19; C12N005-10 WO 200179442 A UPAB: 20040112 NOVELTY - An albumin fusion protein (I) comprising a therapeutic protein: X and (a fragment or variant of) albumin comprising a

fully defined sequence (S18) of 585 amino acids as given in the specification, (where the fragment or variant has albumin or

therapeutic protein: X activity) is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a kit comprising a composition containing (I);
- (2) treating a disease or disorder (that is modulated by therapeutic protein: X or its fragment or variant) comprising administering (I);
- (3) extending the shelf life of therapeutic protein: X comprising fusing therapeutic protein: X or its fragment or variant to albumin or its fragment or variant, sufficient to extend the shelf life of therapeutic protein: X compared to the shelf life of therapeutic protein: X in an unfused state;
- (4) a nucleic acid molecule (II) comprising a polynucleotide sequence encoding (I);
 - (5) a vector comprising (II); and
 - (6) a host cell comprising (II).

ACTIVITY - Cytostatic; anorectic; immunosuppressive; antidiabetic; antirheumatic; antiarthritic; psoriatic. No supporting data is given.

MECHANISM OF ACTION - None given.

USE - **Albumin** fusion proteins are stabilized therapeutic proteins e.g. antibodies to C5, C242 and CD80 useful for treating various diseases and disorders such as non-Hodgkin's lymphoma, cancer, obesity, transplant rejection, type I diabetes mellitus, rheumatoid arthritis and psoriasis.

ADVANTAGE - Fusing albumin to therapeutic proteins stabilizes the therapeutic protein, extends the shelf life and retains the in vitro or in vivo biological activity. It also reduces the need to formulate protein solutions with large excesses of carrier proteins to prevent loss of therapeutic proteins due to factors such as binding to the container. The fusion proteins are easily dispensed with a simple formulation requiring minimal post storage manipulation.

The fusion of therapeutic proteins to **albumin** confers stability in aqueous or other solution. A solution of 200 microgram/ml of human **albumin** (HA)-human growth hormone (hGH) was prepared in tissue culture media containing 5% horse serum and the solution incubated at 37 degrees C starting at time zero. A sample was removed and tested for its biological activity in the Nb2 cell assay at 2 ng/ml final concentration. The biological activity of HA-gHG remained essentially intact after 5 weeks of incubation at 37 degrees C. The recombinant hGH used as control lost its biological activity in the first week of the experiment.

Dwg.0/20

FS' CPI

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TECH

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FA AB; DCN

CPI: B04-B04D4; B04-E02F; B04-E03A; B04-E08; B04-F0100E; B04-G01; B04-N02B0E; B04-P0100E; B11-C07A; B12-K04A; B14-C09B; B14-E12; B14-G02C; B14-H01; B14-N17C; B14-S04; D05-H11; D05-H12A; D05-H12C; D05-H12E; D05-H14; D05-H16; D05-H17C; D05-H17C1 UPTX: 20011129

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Protein: The albumin activity is the ability to prolong the shelf life of the therapeutic protein: X compared to the shelf life of therapeutic protein: X in the unfused state. (I) has a greater shelf life than the therapeutic protein: X in the unfused state. The in vitro or in vivo biological activity of (I) is greater than the in vitro or in vivo activity of therapeutic protein: X or its fragment or variant in an unfused state. (I) comprises 2 therapeutic protein: X or their fragments or variants, which are different from each other. Therapeutic protein: X or its fragment or variant is separated from the albumin or its fragment or variant by a linker. (I) comprises a therapeutic protein: X or its fragment or variant I-inserted into an albumin comprising amino acids 54-61, 76-89, 92-100, 170-176, 247-252, 266-277, 280-288, 362-368, 439-447, 462-475, 478-486 or 560-566 of S18. (I) further comprises a secretion leader sequence. (I) has the formula: R1-L-R2; R2-L-R1; or R1-L-R2-L-R1, where:

R1 = therapeutic protein: X or its fragment or variant;
L = peptide linker; and

R2 = albumin comprising S18.

ABEX

وروايت

(I) is non-glycosylated and expressed in a glycosylation and protease deficient yeast cell. Alternatively (I) is expressed in a mammalian cell in culture.

Preferred Method: The disease or disorder comprises indication: Y. Preparation: (I) are prepared by standard recombinant techniques. UPTX: 20011129

WIDER DISCLOSURE - Also disclosed as new are:

- (1) transgenic organisms modified to contain (II) to express (I);
- (2) antibodies that bind to a therapeutic protein;
- (3) generating antibodies that bind to a therapeutic protein;
- (4) polynucleotides encoding the antibody;
- (5) diagnosing a disorder comprising assaying the expression of the therapeutic protein in cells or body fluid of an individual using antibodies specific to the therapeutic protein and comparing the level of gene expression with a standard gene expression level, where an increase or decrease in the assayed gene expression level is indicative of a particular disorder; and
- (6) a diagnostic kit for use in screening serum containing antigens of a therapeutic protein comprising an antibody immunoreactive with the antigen.

ADMINISTRATION - 0.1-100 mg/kg of body weight, preferably 1-10 mg/kg of body weight of antibodies are administered by standard routes.

EXAMPLE - Preparation of human albumin fusion proteins was as follows. The cDNA for interferon (IFN) alpha was isolated from cDNA libraries by reverse transcription-polymerase chain reaction (PCR) and by PCR using a series of overlapping synthetic oligonucleotides primers using standard methods. The cDNA was tailored at the 5' and 3' ends to generate restriction sites so that oligonucleotide linkers could be used to clone the cDNA into a vector containing the cDNA for human albumin (HA). This could be at the N or C terminus of the HA sequence with(out) use of a spacer sequence. The IFN alpha cDNA was cloned into a vector such as pPPC0005 from which the complete expression cassette was excised and inserted into the plasmid pSAC35 to allow the expression of the albumin fusion protein in yeast. The albumin fusion protein was collected and purified from the media and tested for its biological activity.

- L88 ANSWER 5 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN
- AN **2001-602931** [68] WPIX
- CR 2001-611723 [70]; 2001-616754 [71]; 2001-616755 [71]; 2001-616756 [71]; 2002-010886 [01]; 2002-179329 [23]; 2003-810996 [76]; 2004-033644 [03]

DNC C2001-178694

- TI Albumin fusion proteins comprising a therapeutic protein and albumin, useful in the treating metastatic renal cell carcinoma, metastatic melanoma, malignant melanoma, renal cell carcinoma, HIV (human immunodeficiency virus) or infection.
- DC B04 D16
- IN PRIOR, C P; ROSEN, C A; SADEGHI, H; TURNER, A J
- PA (HUMA-N) HUMAN GENOME SCI INC; (PRIN-N) PRINCIPIA PHARM CORP; (PRIO-I) PRIOR C P; (ROSE-I) ROSEN C A; (SADE-I) SADEGHI H; (TURN-I) TURNER A J CYC 96
- PI WO 2001079258 A1 20011025 (200168)* EN 325p C07K001-00
 - RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW
 - W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

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AU 2001059066 A 20011030 (200219)
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                  A1 20030115 (200313) EN
                                                    C07K001-00
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI TR
     US 2003171267 A1 20030911 (200367)
                                                     A61K038-38
     JP 2003530838 W 20031021 (200373) 430p
                                                    C12N015-09
    WO 2001079258 A1 WO 2001-US12008 20010412; AU 2001059066 A AU 2001-59066
ADT
     20010412; EP 1274720 A1 EP 2001-932549 20010412, WO 2001-US12008 20010412;
     US 2003171267 A1 Provisional US 2000-229358P 20000412, Provisional US
     2000-199384P 20000425, Provisional US 2000-256931P 20001221, US
     2001-833117 20010412; JP 2003530838 W JP 2001-576855 20010412, WO
     2001-US12008 20010412
    AU 2001059066 A Based on WO 2001079258; EP 1274720 Al Based on WO
     2001079258; JP 2003530838 W Based on WO 2001079258
PRAI US 2000-256931P 20001221; US 2000-229358P 20000412; US 2000-199384P
     20000425; US 2001-833117
                                20010412
     ICM A61K038-38; C07K001-00; C12N015-09
         A01N037-18; A61K035-12; A61K035-76; A61K038-00; A61K038-21;
         A61K038-22; A61K038-23; A61K038-27; A61K047-48; A61K048-00;
          A61P001-04; A61P003-10; A61P003-14; A61P005-10; A61P009-10;
          A61P015-08; A61P017-00; A61P017-02; A61P017-06; A61P017-14;
         A61P019-00; A61P019-02; A61P019-08; A61P019-10; A61P021-00;
         A61P025-00; A61P025-02; A61P025-28; A61P029-00; A61P031-14;
         A61P031-18; A61P031-20; A61P035-00; A61P035-02; A61P035-04;
          A61P037-00; A61P037-06; C07K014-55; C07K014-565; C07K014-585;
          C07K014-60; C07K014-62; C07K014-635; C07K014-76; C07K014-765;
          C07K019-00; C12N001-19; C12N005-10
AΒ
    WO 200179258 A UPAB: 20040112
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NOVELTY - Albumin fusion proteins (P1) comprising a therapeutic protein (T1) (or its fragment or variant having the activity of T1) and albumin comprising the 585 amino acid sequence (I) defined in the specification (or its fragment or variant having albumin activity), are new.

 ${\tt DETAILED}$ <code>DESCRIPTION</code> - <code>INDEPENDENT</code> <code>CLAIMS</code> are also included for the following:

- (1) a kit comprising a composition containing P1;
- (2) a method of treating a disease or disorder, preferably modulated by T1, in a patient, comprising administering P1;
- (3) a method of extending the shelf-life of T1, comprising fusing T1 or its fragment or variant, to **albumin** or its fragment or variant, where the shelf-life of T1 or its fragment or variant as part of a fused protein is extended when compared to T1 or its fragment or variant in an unfused state;
 - (4) a nucleic acid (N1) comprising a nucleotide sequence encoding P1;
 - (5) a vector comprising N1; and
 - (6) a host cell comprising N1.

- -

ACTIVITY - Cytostatic; antiviral; antiinflammatory; antileukemic; antiarthritic; antirheumatic; immunosuppressive; antidiabetic; cardiant; nootropic; neuroprotective; antimicrobial; vulnerary.

To test whether sympathetic neuronal cell viability is supported by an albumin fusion protein, the chicken embryo neuronal survival assay (Senaldi, et al., Proc. Natl. Acad., Sci., U.S.A, 96:11458-63 (1998)). Briefly, motor and sympathetic neurons were isolated from chicken embryos, resuspended in L15 medium (with 10% fetal calf serum (FCS), glucose, sodium selenite, progesterone, conalbumin, putrescine and insulin) and Dulbecco's modified Eagles medium (with 10% FCS, glutamine, penicillin, and 25 mM Hepes buffer (pH 7.2)), respectively and incubated at 37 degrees Centigrade in 5% carbon-dioxide in the presence of different concentrations of the purified fusion protein, as well as negative control lacking any cytokine, After 3 days, neuronal survival was determined by evaluation of cellular morphology, and through the use of the colorimetric assay of Mosmann (Mosmann, T., J. Immunol., Methods, 65:55-63 (1983)). Enhanced neuronal cell viability as compared to the

controls lacking cytokine is indicative of the ability of the albumin fusion protein to enhance the survival of neuronal cells. MECHANISM OF ACTION - Gene therapy.

USE - When the therapeutic protein, or its fragment or variant is IL-2, P1 is used to treat metastatic renal cell carcinoma, metastatic melanoma, malignant melanoma, renal cell carcinoma, HIV (human immunodeficiency virus) infection, inflammatory bowel disorder, Kaposi's sarcoma, leukemia, multiple sclerosis, rheumatoid arthritis, transplant rejection, type 1 diabetes mellitus, lung cancer, acute myeloid leukemia, hepatitis C, non-hodgkin's lymphoma or ovarian cancer (claimed).

The albumin fusion proteins are also useful in the treatment, prevention, diagnosis, and/or detection of diseases, disorders such as immune system disorders (e.g. transplant rejection), blood related disorders (e.g. myocardial infarction), hyperproliferative disorders (e.g. childhood acute myeloid leukemia), renal disorders (e.g. glomerulonephritis), cardiovascular disorders (e.g. arrhythmias), respiratory disorders (e.g. non-allergic rhinitis), neurological diseases (e.g. Alzheimer's disease), endocrine disorders (e.g. pheocytochroma), reproductive system disorders (e.g. syphilis), infectious diseases (e.g. measles), gastrointestinal disorders (e.g. irritable bowel syndrome) and wound healing.

Dwg.0/14

FS CPI

17.

-F.

AB; DCN FΑ

CPI: B04-C01; B04-E02F; B04-E08; B04-F0100E; B04-F1100E; MC

B04-H05; B04-H06; B04-J04; B04-N0200E;

B04-N02A0E; B14-A02B1; B14-C09B; B14-D01; B14-E10C; B14-F01; B14-F02; B14-G02; B14-H01; B14-J01; B14-K01; B14-N10; B14-N12;

B14-N14; B14-N17B; B14-S01; B14-S03; B14-S04; **D05-H12B2**;

D05-H12E; D05-H14

UPTX: 20011121 TECH

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Fusion Protein: The albumin activity is the ability to prolong the shelf-life of T1 compared to the shelf-life of T1 in an unfused state. The albumin fragment or variant comprises amino acids 1-387 of (I). T1 comprises interleukin 2 (IL-2). The T1 fragment or variant has T cell proliferative activity or T cell activation activity. Tl or its fragment or variant, comprises a protein selected from calcitonin, growth hormone releasing factor, IL-2 fusion protein, insulin-like growth factor-1, interferon beta or parathyroid hormone. T1 or its fragment or variant is fused to the C-terminal of the albumin or the C-terminus of the fragment or variant of albumin. Alternatively, T1 or its fragment or variant is fused to the N-terminal of the albumin or the N-terminus of the fragment or variant of albumin. Alternatively, T1 or its fragment or variant is fused to the N-terminus and C-terminus of the albumin, or the N-terminus and C-terminus of the fragment or variant of albumin. P1 comprises a first T1 or its fragment or variant, and a second T1 or its fragment or variant, where the first T1 is different from the second T1. Tl or its fragment or variant is separated from the albumin or the fragment or variant of albumin by a linker. Preferably, P1 is of the formula (S1), (S2) or (S3). R1-L-R2 (S1); R2-L-R1 (S2); or R1-L-R2-L-R1 (S3). R1 = is T1 or its fragment or variant;

L = is a peptide linker; and

R2 = is albumin comprising the sequence of (I), or its fragment

The shelf-life of the albumin fusion protein is greater than the shelf-life of T1 or its fragment or variant in an unfused state. The in vitro or in vivo biological activity of Tl or its fragment or variant, fused to **albumin** or its fragment or variant, is greater than the in vitro or in vivo, respectively, biological activity of T1 or its fragment or variant, in an unfused state. Alternatively, P1 comprises T1 or its fragment or variant, inserted into an **albumin** comprising the sequence of (I) or its fragment or variant. Preferably, the **albumin** comprises residues 54-61, 76-89, 92-100, 170-176, 247-252, 266-277, 280-288, 362-368, 439-447, 462-475, 478-486, or 560-566 of (I). The portion of **albumin** is sufficient to prolong the shelf-life and in vitro and in vivo biological activity of T1 or its fragment or variant, as compared to the shelf-life and in vitro and in vivo biological activity of T1 or its fragment or variant, in an unfused state. P1 is non-glycosylated and expressed in yeast which is glycosylation deficient. The yeast may also be protease deficient. Alternatively, P1 is

expressed by a mammalian cell in culture. Pl further comprises a secretion

leader sequence.
ABEX UPTX: 20011121

ADMINISTRATION - The **albumin** fusion proteins can be administered orally, rectally, parenterally, intracisternally, intravaginally, intraperitoneally, topically, bucally, or as an oral or nasal spray. The dosage is 1 microgram/kg/day to 10 mg/kg/day, preferably 0.01 to 1, mg/kd/day. If given continuously, the **albumin** fusion protein is typically administered at a dose rate of 1-50 micrograms/kg/hour, either by 1-4 injections per day or by continuous subcutaneous infusions.

EXAMPLE - The cDNA for the growth factor of interest such as interferon growth factor 1 (IGF-1) can be isolated using a variety of means including but not exclusively, from cDNA libraries, by reverse transcriptasepolymerase chain reaction (PCR) and by PCR using a series of overlapping synthetic oligonucleotide primers, all using standard methods (see GenBank Acc. Number NP-000609). The cDNA can be tailored at the 5' and 3' ends to generate restriction sites, such that the oligonucleotide linkers can be used, for cloning of the cDNA into a vector containing the cDNA for human serum albumin (HA). This can be a the N or C-terminus with or without the use of a spacer sequence. The growth factor cDNA was cloned into a vector such as pPPC0005, pScCHSA, pScNHSA or pC4:HSA from which the complete expression cassette is then excised and inserted into the plasmid pSAC35 to allow the expression of the albumin fusion protein in yeast. The albumin fusion protein secreted from the yeast can then be collected and purified from the media and tested for its biological activity. For expression in mammalian cell lines a similar procedure is adopted except that the expression cassette used employs a mammalian promoter, leader sequence and terminator. This expression cassette is then excised and inserted into a plasmid suitable for the transfection of mammalian cell lines.

L88 ANSWER 6 OF 6 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 1996-300388 [30] WPIX

DNC C1996-095415

TI New chimeric proteins for treatment of septic shock, psoriasis, cancers etc. - comprise cytokine bonded to polypeptide which is enzymatically inactive in humans, increases half-life and prevents cytokine(s) from crossing blood brain barrier.

DC B04

ور والمحقود

IN STEELE, A; STROM, T B; ZHENG, X; ZHENG, X X

PA (BETH-N) BETH ISRAEL HOSPITAL ASSOC

CYC 20

PI WO 9618412 A1 19960620 (199630)* EN 58p A61K038-19 RW: AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT SE W: CA JP

EP 793504 A1 19970910 (199741) EN A61K038-19

R: CH DE FR GB IT LI SE

JP 11501506 W 19990209 (199916) 49p C12N015-09

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A61K038-20
                   B1 20020611 (200244)
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                  B1 20020625 (200246)
                                                     C07K014-54
     US 6410008
     US 2002173628 A1 20021121 (200279)
                                                     A61K038-52
     US 2003026778. A1 20030206 (200318)
                                                     A61K038-20
     WO 9618412 A1 WO 1995-US16046 19951212; EP 793504 A1 EP·1995-943058
     19951212, WO 1995-US16046 19951212; JP 11501506 W WO 1995-US16046
     19951212, JP 1996-519191 19951212; US 6403077 B1 CIP of US 1994-355502
     19941212, Cont of US 1995-431535 19950428, US 1997-968905 19971106; US
     6410008 B1 US 1994-355502 19941212; US 2002173628 A1 Cont of US
     1994-355502 19941212, US 2002-145481 20020514; US 2003026778 A1 CIP of US
     1994-355502 19941212, Cont of US 1997-968905 19971106, US 2002-145517
     20020514
     EP 793504 Al Based on WO 9618412; JP 11501506 W Based on WO 9618412; US
     2002173628 A1 Cont of US 6410008; US 2003026778 A1 Cont of US 6403077, CIP
     of US 6410008
PRAI US 1995-431535
                      19950428; US 1994-355502
                                                 19941212; US 1997-968905
     19971106; US 2002-145481
                              20020514; US 2002-145517
     2.Jnl.Ref; US 5231012
     ICM A61K038-19; A61K038-20; A61K038-52; C07K014-54; C12N015-09
         A61K038-00; A61K038-21; A61K038-38; A61K039-395;
          C07K014-52; C07K014-525; C07K014-53; C07K014-535;
          C07K014-545; C07K014-55; C07K014-555; C07K014-76;
          C07K014-765; C07K016-18; C07K016-46; C07K019-00;
          C12N009-10; C12N015-02; C12N015-24; C12P021-02
AΒ
          9618412 A UPAB: 19960731
     Chimeric protein comprises a cytokine bonded to a polypeptide which is
     enzymatically inactive in humans and which increases the circulating
     half-life of the cytokine in vivo by a factor of 1.
           Also claimed is the use of interleukin-10 (IL-10)/Fc in the preparation
     of a medicament for inhibiting granuloma formation in a patient.
          USE - The chimeric proteins can be used to treat conditions for which
     the corresp. cytokines are used, e.g. septic shock, granulomatous
     disorders (e.g. schistosomiasis), multiple sclerosis, psoriasis,
     rheumatoid arthritis, cancers and virus infections. Chimeric proteins
     including a lytic Fc region can also be used to deplete patients of
     suppressor lymphocytes and to treat chronic infections such as those
     associated with suppression of the immune system.
          ADVANTAGE - The enzymatically inactive polypeptides extend the
     circulating half-life of the cytokines in vivo by a factor of 10
     (claimed). In addition, they can prevent the cytokines from crossing the
     blood brain barrier and causing adverse side effects.
     Dwg.0/15
     CPI
FS
FΑ
     AB
     CPI: B04-B04; B04-G01; B04-H02; B04-H04A; B04-H04C; B04-H08;
MC
          B04-N02; B14-A01; B14-C09B; B14-N17C; B14-S01; B14-S06
=> => d his
     (FILE 'HOME' ENTERED AT 15:22:31 ON 02 FEB 2004)
                SET COST OFF
     FILE 'HCAPLUS' ENTERED AT 15:22:50 ON 02 FEB 2004
                E ALBUMIN/CT
            753 S E3
L1
            132 S E11
L2
                E E47+ALL
          80101 S E2+NT
L3
                E E33+ALL
L4
            566 S E3, E2
          25218 S E2+NT
L5
         157881 S ?ALBUMIN?
L6
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مه والمتشد

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181833 S L1-L6
L7
            2969 S BDNF OR BD NF
L8
            2881 S BRAIN DERIVED NEUROTROPHIC FACTOR
L9
            2883 S (BD OR BRAIN DERIVED) () (NF OR NEUROTROPHIC FACTOR)
L10
                 E NEUROTROPHIC FACTOR/CT
L11
            141 S E10
            2554 S E26
                 E E25+ALL
             789 S E3-E5 AND BRAIN DERIVED
L13
L14
            679 S E12,E13
L15
            3242 S E2+NT (L) BRAIN DERIVED
L16
              64 S L7 AND L8-L15
           19234 S INTERFERONALPHA OR ALPHAINTERFERON OR INTERFERONBETA OR BETAI
L17
                 E INTERFERON/CT
· L18
             302 S E3-E19
L19
           18390 S E85-E101
                 E INTERFERONS/CT
                 E E3+ALL
           18391 S E7, E6 (L) (ALPHA OR BETA)
L21
             546 S L7 AND L17-L20
            2340 S TIMP()(I OR 1)
L22
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L23
               1 S 140208-24-8
      FILE 'HCAPLUS' ENTERED AT 15:30:37 ON 02 FEB 2004
L24
            2026 S L23
L25
             859 S TISSUE INHIBITOR (1W) METALLOPROTEINASE 1
L26
              27 S METALLOPROTEINASE INHIBITOR 1
L27
             651 S TIMP1
             12 S FIBROBLAST COLLAGENASE INHIBITOR
L28
L29
             91 S L7 AND L22, L24-L28
L30
             678 S L16, L21, L29
            9815 S IFNALPHA OR IFNBETA OR ALPHAIFN OR BETAIFN OR IFN(A) (ALPHA OR
L31
L32 ·
             119 S L7 AND L31
             700 S L30, L32
L33
             62 S L33 AND (FUSION OR FUSE OR FUSED OR FUSES OR FUSING)
L34
L35
             167 S L33 AND RECOMBIN?
L36
              44 S L33 AND CHIMER?
L37
             202 S L34-L36
                 E ROSEN C/AU
 L38
              27 S E3, E4
                 E ROSEN CRAIG/AU
             625 S E3-E5
                 E HASELTINE W/AU
L40
             302 S E3, E4, E7-E10
              10 S L33 AND L38-L40
L41
                 E HUMAN GENOME SCI/PA, CS
             975 S E5-E37
L43
              13 S L33 AND L42
              13 S L41, L43
L44
              13 S L44 AND L37
L45
L46
               9 S L45 AND (SHELFLIFE OR SHELF LIFE)
L47
               4 S L45 NOT L46
                 SEL DN AN 1 4
               2 S L47 NOT E1-E6
              11 S L46, L48
L49
                 SEL RN
                 DEL SEL
                 E FUSION PROTEIN/CT
 L50 ·
           11933 S E9
                 E E9+ALL
            3795 S E3, E4
 L51
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5 S L51 AND L33
L52
             29 S L50 AND L33
L53
             34 S L49, L52, L53
L54
L55
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             7 S L54 NOT L55
L56
            159 S L37 AND ALBUMIN
L57
            132 S L57 NOT L43-L49, L52-L56
L58
              6 S L58 AND L16
L59
             7 S L58 AND L29 '
L60
            121 S L58 NOT L59, L60
L61
             96 S L61 AND (PD<=20000412 OR PRD<=20000412 OR AD<=20000412)
L62
                SEL DN AN 9 12 13 24 29 31 35 39 44 47 55 58 72 74 83 85 92 93
             18 S L62 AND E1-E54
L63
             29 S L49, L63 AND L1-L22, L24-L63
L64
             29 S L64 AND ?ALBUMIN?
L65
             29 S L64 AND (INF? OR INTERFERON OR TIMP? OR NEUROTROPHIC?)
L66
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     FILE 'WPIX' ENTERED AT 16:01:33 ON 02 FEB 2004
           9861 S L6/BIX
L67
L68
           318 S L8/BIX OR L9/BIX OR L10/BIX
           1564 S L17/BIX OR LL31/BIX
L69
            80 S L22/BIX OR L25/BIX OR L26/BIX OR L27/BIX OR L28/BIX
L70
L71
            124 S L67 AND L68-L70
          11209 S ?ALBUMEN?/BIX OR L67
L72
            513 S (A61K038-38 OR C07K014-76 OR C07K014-765 OR C12N015-14)/IC,IC
L73
L74
          11377 S L72,L73
          2983 S V275/M0, M1, M2, M3, M4, M5, M6 OR (B02-V03 OR C02-V03 OR B04-H05A
L75
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L76
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L77
            111 S L74 AND L76
L78
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L:79
L80
            311 S L77-L79
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L81
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L82
L83
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L84
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L85
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L86
L87
              6 S L81, L86
L88
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     FILE 'HCAPLUS' ENTERED AT 16:25:16 ON 02 FEB 2004
     FILE 'REGISTRY' ENTERED AT 16:26:59 ON 02 FEB 2004
L89
             1 S 507485-69-0
L90
              1 S 472960-22-8
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                 CA/CAplus records now contain indexing from 1907 to the
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         SEP 09
                 present
                 INPADOC: Legal Status data reloaded
NEWS
         DEC 08
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NEWS
      5
         SEP 29
         OCT 10
NEWS
     6
                 PCTFULL: Two new display fields added
                 BIOSIS file reloaded and enhanced
NEWS
      7
         OCT 21
                 BIOSIS file segment of TOXCENTER reloaded and enhanced
         OCT 28
NEWS
     8
                 MSDS-CCOHS file reloaded
         NOV 24
NEWS
     9
                 CABA reloaded with left truncation
         DEC 08
NEWS 10
         DEC 08
                 IMS file names changed
NEWS 11
                 Experimental property data collected by CAS now available
NEWS 12
         DEC 09
                 in REGISTRY
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NEWS 13
         DEC 09
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NEWS 14
         DEC 17
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NEWS 15
         DEC 18
                 CROPU no longer updated; subscriber discount no longer
         DEC 19
NEWS 16
                 available
                 Additional INPI reactions and pre-1907 documents added to CAS
         DEC 22
NEWS 17
                 databases
                 IFIPAT/IFIUDB/IFICDB reloaded with new data and search fields
         DEC 22
NEWS 18
         DEC 22
                 ABI-INFORM now available on STN
NEWS 19
                 Source of Registration (SR) information in REGISTRY updated
         JAN 27
NEWS 20
                 and searchable
                 A new search aid, the Company Name Thesaurus, available in
         JAN 27
NEWS 21
                 CA/CAplus
                 German (DE) application and patent publication number format
NEWS 22
         FEB 05
                 changes
              DECEMBER 28 CURRENT WINDOWS VERSION IS V7.00, CURRENT
NEWS EXPRESS
              MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP),
              AND CURRENT DISCOVER FILE IS DATED 23 SEPTEMBER 2003
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FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004

=> file medline, uspatful, dgene, embase, wpids

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'MEDLINE' ENTERED AT 13:52:47 ON 06 FEB 2004

FILE 'USPATFULL' ENTERED AT 13:52:47 ON 06 FEB 2004 CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'DGENE' ENTERED AT 13:52:47 ON 06 FEB 2004 COPYRIGHT (C) 2004 THOMSON DERWENT

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FILE 'WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004 COPYRIGHT (C) 2004 THOMSON DERWENT

=> s albumin fusion proteins

2835 ALBUMIN FUSION PROTEINS

=> s cerebus protein

1 CEREBUS PROTEIN

=> s 11 and 12

0 L1 AND L2

=> s (cerebus protein) and albumin

0 (CEREBUS PROTEIN) AND ALBUMIN

=> s 12 and fusion

0 L2 AND FUSION

=> d 12 ti abs ibib tot

ANSWER 1 OF 1 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L₂

TI Human and murine cerebus-like proteins - used for treating tissue defects and degenerative nerve conditions.

NΑ 1999-106054 [09] WPIDS

CR 2003-298696 [29]

9901553 A UPAB: 20030505 AΒ

A novel isolated DNA sequence comprises a DNA sequence selected from: (a) nucleotides beginning at # 1, 52, 55, 58, 61, 64, 67, 70, 73, 121, 256, 259, 262, 265, 268, 171, or 484 and ending at # 723 or 801 of the 804 bp DNA sequence given in the specification; and (b) sequences which hybridise to (a) under stringent hybridisation conditions and encode a protein which exhibits cerebus activity. Also claimed are: (1) an isolated DNA sequence comprising nucleotides encoding amino acids beginning at #1, 18 to 25, 41, 85 to 91 or 152, and ending at #241 or 267 of the 267 amino acid sequence given in the specification; (2) a vector comprising either of the above DNA molecules in operative association with an expression control sequence; (3) an isolated DNA molecule comprising nucleotides 268-801 of the 272 amino acid sequence given in the specification (sic), or naturally occurring allelic sequences of it; (4) a vector comprising the DNA of (4) in operative association with an expression control sequence; (5) an isolated DNA molecule encoding mammalian cerebus protein , comprising nucleotides 268-801 of the 804 bp DNA sequence given in the specification; (6) a vector comprising the DNA of (5) in operative

association with an expression control sequence; (7) a host cell

transformed with the vector of (2), (4) or (6); (8) a purified mammalian cerebus protein comprising the 267 amino acid sequence given in the specification; (9) a purified mammalian cerebus protein comprising residues 90-267 of the 272 amino acid sequence given in the specification; and (10) antibodies to the cerebus protein of (8) or (9).

USE - The host cell of (7) can be used to produce the mammalian cerebus proteins (claimed). Compositions containing the protein can be used in the formation of neurons and related neural cells and tissues, such as Schwann cells, glial cells, and astrocytes, as well as liver, pancreas, lung, heart, kidney, spleen, stomach, and cardiac tissue and cells. They may also be used to treat precursor or stem cells. The compositions can also be used for treating tissue defects, and healing and maintenance of various types of tissues and wounds. The mammalian cerebus protein containing compositions may also be used to treat or prevent degenerate nerve conditions such as Parkinson's disease, Alzheimer's disease, and Lou Gehrig's disease. They can also be used to treat osteoporosis, rheumatoid arthritis, osteoarthritis, and other abnormalities of connective tissue, or of other organs or tissues.

ACCESSION NUMBER:

1999-106054 [09] WPIDS

CROSS REFERENCE:

2003-298696 [29]

DOC. NO. CPI:

C1999-031758

TITLE:

Human and murine cerebus-like proteins - used for treating tissue defects and degenerative nerve

conditions.

DERWENT CLASS:

B04 D16

INVENTOR(S):

DEROBERTIS, E M; FOLLETTIE, M

PATENT ASSIGNEE(S):

(GEMY) GENETICS INST INC; (REGC) UNIV CALIFORNIA

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9901553 A1 19990114 (199909)* EN 50

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

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AU 9878140 A 19990125 (199923)

US 5935852 A 19990810 (199938)

EP 1012278 A1 20000628 (200035) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

MX 2000000242 A1 20010601 (200235)

JP 2002511762 W 20020416 (200242)

AU 749031 B 20020620 (200252)

APPLICATION DETAILS:

PAT	ENT NO K	IND	API	PLICATION	DATE
WO	9901553	A1	wo	1998-US11462	19980603
AU	9878140	A	ΑU	1998-78140	19980603
US	5935852	A	US	1997-887997	19970703
ΕP	1012278	A1	ΕP	1998-926263	19980603
			WO	1998-US11462	19980603
MX	2000000242	A1	MX	2000-242	20000105
JP	2002511762	W	WO	1998-US11462	19980603
			JΡ	1999-507147	19980603
ΑU	749031	В	ΑU	1998-78140	19980603

FILING DETAILS:

PATENT NO KIND PATENT NO							
AU 9878140 EP 1012278 JP 2002511762 AU 749031	A1 W	Based on Based on Based on Previous Publ Based on	WO WO LA	9901553 9901553 9901553 9878140 9901553			

PRIORITY APPLN. INFO: US 1997-887997 19970703

=> file biosis

SINCE FILE TOTAL COST IN U.S. DOLLARS ENTRY SESSION

20.32 20.53 FULL ESTIMATED COST

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FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 4 February 2004 (20040204/ED)

FILE RELOADED: 19 October 2003.

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

2835 S ALBUMIN FUSION PROTEINS LI

1 S CEREBUS PROTEIN 1.2

0 S L1 AND L2 L3

0 S (CEREBUS PROTEIN) AND ALBUMIN 1.4

0 S L2 AND FUSION L5

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

=> s 12

0 CEREBUS

1361492 PROTEIN

O CEREBUS PROTEIN L6

(CEREBUS (W) PROTEIN)

=> file medline, uspatful, dgene, embase, wpids, biosis, japio, fsta, jicst TOTAL SINCE FILE

COST IN U.S. DOLLARS

FULL ESTIMATED COST

SESSION

21.38

ENTRY 0.85

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FILE 'USPATFULL' ENTERED AT 14:00:26 ON 06 FEB 2004 CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE 'FSTA' ENTERED AT 14:00:26 ON 06 FEB 2004 COPYRIGHT (C) 2004 International Food Information Service

FILE 'JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004 COPYRIGHT (C) 2004 Japan Science and Technology Agency (JST)

=> s 12

L7 1 L2

=> d 17 ti abs ibib tot

L7 ANSWER 1 OF 1 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN TI Human and murine cerebus-like proteins - used for treating tissue defects

and degenerative nerve conditions. AN 1999-106054 [09] WPIDS

CR 2003-298696 [29]

AB WO 9901553 A UPAB: 20030505

A novel isolated DNA sequence comprises a DNA sequence selected from: (a) nucleotides beginning at # 1, 52, 55, 58, 61, 64, 67, 70, 73, 121, 256, 259, 262, 265, 268, 171, or 484 and ending at # 723 or 801 of the 804 bp DNA sequence given in the specification; and (b) sequences which hybridise to (a) under stringent hybridisation conditions and encode a protein which exhibits cerebus activity. Also claimed are: (1) an isolated DNA sequence comprising nucleotides encoding amino acids beginning at #1, 18 to 25, 41, 85 to 91 or 152, and ending at #241 or 267 of the 267 amino acid sequence given in the specification; (2) a vector comprising either of the above DNA molecules in operative association with an expression control sequence; (3) an isolated DNA molecule comprising nucleotides 268-801 of the 272 amino acid sequence given in the specification (sic), or naturally occurring allelic sequences of it; (4) a vector comprising the DNA of (4) in operative association with an expression control sequence; (5) an isolated DNA molecule encoding mammalian cerebus protein , comprising nucleotides 268-801 of the 804 bp DNA sequence given in the specification; (6) a vector comprising the DNA of (5) in operative association with an expression control sequence; (7) a host cell transformed with the vector of (2), (4) or (6); (8) a purified mammalian cerebus protein comprising the 267 amino acid sequence given in the specification; (9) a purified mammalian cerebus protein comprising residues 90-267 of the 272 amino acid sequence given in the specification; and (10) antibodies to the cerebus protein of (8) or (9).

USE - The host cell of (7) can be used to produce the mammalian cerebus proteins (claimed). Compositions containing the protein can be used in the formation of neurons and related neural cells and tissues, such as Schwann cells, glial cells, and astrocytes, as well as liver, pancreas, lung, heart, kidney, spleen, stomach, and cardiac tissue and cells. They may also be used to treat precursor or stem cells. The compositions can also be used for treating tissue defects, and healing and maintenance of various types of tissues and wounds. The mammalian cerebus protein containing compositions may also be used to treat or prevent degenerate nerve conditions such as Parkinson's disease, Alzheimer's disease, and Lou Gehrig's disease. They can also be used to treat osteoporosis, rheumatoid arthritis, osteoarthritis, and other abnormalities of connective tissue, or of other organs or tissues. Dwg.0/0

ACCESSION NUMBER:

1999-106054 [09] WPIDS

CROSS REFERENCE:

2003-298696 [29]

DOC. NO. CPI:

C1999-031758

TITLE:

Human and murine cerebus-like proteins - used for

treating tissue defects and degenerative nerve

conditions.

DERWENT CLASS:

B04 D16

INVENTOR(S):

DEROBERTIS, E M; FOLLETTIE, M

PATENT ASSIGNEE(S):

(GEMY) GENETICS INST INC; (REGC) UNIV CALIFORNIA

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA PG	ţ
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A1 19990114 (199909) * EN WO 9901553 50

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG

57

UZ VN YU ZW

AU 9878140 A 19990125 (199923) US 5935852 A 19990810 (199938)

EP 1012278 A1 20000628 (200035) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

MX 2000000242 A1 20010601 (200235)

JP 2002511762 W 20020416 (200242) AU 749031 B 20020620 (200252)

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 9901553	A1	WO 1998-US11462	19980603
AU 9878140	A	AU 1998-78140	19980603
US 5935852	A	US 1997-887997	19970703
EP 1012278	A1	EP 1998-926263	19980603
		WO 1998-US11462	19980603
MX 2000000242	A1	MX 2000-242	20000105
JP 2002511762	W	WO 1998-US11462	19980603
		JP 1999-507147	19980603
AU 749031	В	AU 1998-78140	19980603

FILING DETAILS:

PAT	TENT NO K	IND			PA	TENT NO
	9878140 1012278		Based on Based on			9901553 9901553
JP	2002511762		Based on		WO	9901553.
AU	749031	В	Previous Based on	Publ.		9878140 9901553

PRIORITY APPLN. INFO: US 1997-887997 19970703

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(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L21 S CEREBUS PROTEIN L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

=> s TIMP-1 or tissue inhibitor metalloproteinase-1

5 FILES SEARCHED...

8080 TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

=> s 18 and 11

L9 5 L8 AND L1

=> d 19 ti abs ibib tot

L9 ANSWER 1 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:
INVENTOR(S):

Albumin fusion proteins Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 2 OF 5 USPATFULL on STN

TI Albumin fusion proteins

AB The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic

acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003219875	A1	20031127	
APPLICATION INFO.:	US 2001-833118	A1	20010412	(9)

PRIORITY INFORMATION:

US 2000-256931P 20001221 (60) US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 29

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 3 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL

TITLE:
INVENTOR(S):

Albumin fusion proteins
Ballance, David J., Berwyn, PA, UNITED STATES
Sleep, Darrell, West Bridgford, UNITED KINGDOM

Prior, Christopher P., Rosemont, PA, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003199043	A1	20031023	
APPLICATION INFO.:	US 2001-832501	A1	20010412	(9)

NUMBER DATE

_____ ____

PRIORITY INFORMATION:

US 2000-256931P

20001221 (60)

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility
APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

60

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 4 OF 5 USPATFULL on STN

TI Albumin fusion proteins

AB The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:244853 USPATFULL Albumin fusion proteins

TITLE: INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003171267	A1	20030911	
APPLICATION INFO.:	US 2001-833117	A1	20010412	(9)

			NUMBER	DATE	
PRIORITY	INFORMATION:		2000-256931P 2000-199384P	20001221 20000425	
		US	2000-229358P	20000412	(60)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 59 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 20 Drawing Page(s)

LINE COUNT: 13208

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L9 ANSWER 5 OF 5 USPATFULL on STN
- TI Albumin fusion proteins
- The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion

proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT. ACCESSION NUMBER: 2003:181414 USPATFULL Albumin fusion proteins TITLE: Rosen, Craig A., Laytonsville, MD, UNITED STATES INVENTOR(S): Haseltine, William A., Washington, DC, UNITED STATES NUMBER KIND DATE _______ US 2003125247 A1 20030703 PATENT INFORMATION: US 2001-833041 A1 20010412 (9) APPLICATION INFO.: NUMBER DATE _____ US 2000-256931P 20001221 (60) PRIORITY INFORMATION: 20000425 (60) US 2000-199384P US 2000-229358P 20000412 (60) DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE: ROCKVILLE, MD, 20850 NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 20 Drawing Page(s) NUMBER OF DRAWINGS: 15235 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. => d his (FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004) FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004 2835 S ALBUMIN FUSION PROTEINS L11 S CEREBUS PROTEIN L20 S L1 AND L2 L30 S (CEREBUS PROTEIN) AND ALBUMIN L40 S L2 AND FUSION L5 FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004 0 S L2 1.6 FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004 1 S L2 L78080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1 L8 5 S L8 AND L1 L9 => s 18 and fusion 378 L8 AND FUSION L10=> s 110 and albumin 221 L10 AND ALBUMIN L11=> s 111 and albumin fragment 5 L11 AND ALBUMIN FRAGMENT => d l12 ti abs ibib tot

L12 ANSWER 1 OF 5 USPATFULL on STN
TI Albumin fusion proteins

AB The present invention encompasses albumin fusion

proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

•	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2004010134	A1	20040115	
APPLICATION INFO.:	US 2001-833245	A1	20010412	(9)
· · · · · · · · · · · · · · · · · · ·				

NUMBER DATE PRIORITY INFORMATION: US 2000-256931P 20001221 (60)

DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 2 OF 5 USPATFULL on STN

Albumin fusion proteins ТT

AB The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

		NUMBER	KIND	DATE	
PATENT INFORMATION:	ບຣ	2003219875	A1	20031127	
APPLICATION INFO.:	US	2001-833118	A1	20010412	(9)

NUMBER DATE PRIORITY INFORMATION: US 2000-256931P 20001221 (60)

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 3 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL

INVENTOR (S):

TITLE:

Albumin fusion proteins
Ballance, David J., Berwyn, PA, UNITED STATES
Sleep, Darrell, West Bridgford, UNITED KINGDOM
Prior, Christopher P., Rosemont, PA, UNITED STATES
Sadeghi, Homayoun, Doylestown, PA, UNITED STATES
Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003199043	A1	20031023	
APPLICATION INFO.:	US 2001-832501	A1	20010412	(9)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 60 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 4 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using

these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:244853 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003171267	A1	20030911	
APPLICATION INFO.:	US 2001-833117	A1	20010412	(9)

DOCUMENT TYPE: FILE SEGMENT: Utility APPLICATION

LEGAL REPRESENTATIVE: HUM

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

59 1

NUMBER OF DRAWINGS:

20 Drawing Page(s)

LINE COUNT:

13208

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 5 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:181414 USPATFULL Albumin fusion proteins

INVENTOR (S):

TITLE:

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

20000425 (60)

20000412 (60)

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003125247 US 2001-833041	A1 A1	20030703 20010412	(9)
	NUMBER	DA'	TE	
PRIORITY INFORMATION:	US 2000-256931P	2000	1221 (60)	

US 2000-199384P

US 2000-229358P

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 29

NUMBER OF DRAWINGS:

20 Drawing Page(s)

LINE COUNT:

15235

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

L8 8080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

L9 5 S L8 AND L1 L10 378 S L8 AND FUSION L11 221 S L10 AND ALBUMIN

L12 5 S L11 AND ALBUMIN FRAGMENT

=> s l11 and shelf-life

L13 9 L11 AND SHELF-LIFE

=> d l13 ti abs ibib tot

L13 ANSWER 1 OF 9 USPATFULL on STN

TI Biospecific contrast agents

Methods and apparatuses for detecting a condition of a sample (including cervical cancers and pre-cancers) through reflectance and/or fluorescence imaging. A sample is obtained. One or more metallic nanoparticles and/or one or more quantum dots are obtained. The one or more metallic nanoparticles and/or one or more quantum dots are coupled to one or more biomarkers of the sample that are associated with the condition. A reflectance and/or fluorescence image of the sample is then taken. The image(s) exhibit characteristic optical scattering from the one or more metallic nanoparticles and/or characteristic fluorescence excitation from the one or more quantum dots to signal the presence of the one or more biomarkers. In this way, the condition can be readily screened or diagnosed.

ACCESSION NUMBER:

2004:31276 USPATFULL

TITLE:

Biospecific contrast agents

INVENTOR(S):

Sokolov, Konstantin, Austin, TX, UNITED STATES Korgel, Brian A., Round Rock, TX, UNITED STATES Ellington, Andrew D., Austin, TX, UNITED STATES Richards-Kortum, Rebecca, Austin, TX, UNITED STATES

NUMBER KIND DATE

PATENT INFORMATION:

US 2004023415 A1

20040205

APPLICATION INFO.:

US 2003-382136

20030305 (10)

NUMBER DATE

Δ1

PRIORITY INFORMATION:

US 2002-361924P

20020305 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

Michael C. Barrett, Esq., FULBRIGHT & JAWORSKI, L.L.P.,

600 Congress Avenue, Suite 2400, Austin, TX, 78701

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 44

NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT:

3948

L13 ANSWER 2 OF 9 USPATFULL on STN

TI Albumin fusion proteins

AB The present invention encompasses albumin fusion

proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:
INVENTOR(S):

Albumin fusion proteins
Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2004010134	A1	20040115	
APPLICATION INFO.:	US 2001-833245	A1	20010412	(9)

NUMBER DATE

US 2000-256931P 20001221 (60) US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

US 2000-229358P
DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

PRIORITY INFORMATION:

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 3 OF 9 USPATFULL on STN

TI Nanoporous particle with a retained target

AB Porous nanostructured materials, such as porous nanostructured liquid and liquid crystalline particles or materials, incorporate a target substantially within the material which selectively binds a chemical of interest which can diffusion within the porous nanostructured material and be bound by the target. The porous nanostructured materials can be dispersed as particles in a medium in which said chemical of interest is located with low turbidity. Markers which detect binding of said

chemical of interest can be maintained in the medium separate and apart from the target, and any active compound (e.g., an enzyme) associated therewith by the porous nanostructured material, such that detectable changes in the marker only result when the active compounds diffuse out of the porous nanostructured materials after the chemical of interest binds to the target.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:330129 USPATFULL

TITLE:

Nanoporous particle with a retained target

INVENTOR (S):

Anderson, David, Colonial Heights, VA, UNITED STATES

NUMBER KIND DATE -----20031218

PATENT INFORMATION:

A1 US 2003232340

APPLICATION INFO .:

US 2002-170214 A1 20020613

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

WHITHAM, CURTIS & CHRISTOFFERSON, P.C., 11491 SUNSET

HILLS ROAD, SUITE 340, RESTON, VA, 20190

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

119 1

NUMBER OF DRAWINGS:

2 Drawing Page(s)

LINE COUNT:

2555

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 4 OF 9 USPATFULL on STN

TΙ Albumin fusion proteins

AB

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL Albumin fusion proteins

TITLE: INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
		-		
PATENT INFORMATION: US	2003219875	A1	20031127	
APPLICATION INFO.: US	2001-833118	A1	20010412	(9)

			NUMBER	DATE	
PRIORITY	INFORMATION:	US	2000-256931P 2000-199384P 2000-229358P	20001221 20000425 20000412	(60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

LEGAL REPRESENTATIVE:

APPLICATION HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

29

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 5 OF 9 USPATFULL on STN

ΤI Albumin fusion proteins

The present invention encompasses albumin fusion AB proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Ballance, David J., Berwyn, PA, UNITED STATES Sleep, Darrell, West Bridgford, UNITED KINGDOM Prior, Christopher P., Rosemont, PA, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003199043 US 2001-832501	A1	20031023 20010412	(9)

NUMBER DATE US 2000-256931P 20001221 (60) PRIORITY INFORMATION: US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

1

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 6 OF 9 USPATFULL on STN

ΤI Albumin fusion proteins

The present invention encompasses albumin fusion AB proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER: 2003:244853 USPATFULL TITLE: Albumin fusion proteins

Rosen, Craig A., Laytonsville, MD, UNITED STATES INVENTOR(S):

Sadeghi, Homayoun, Doylestown, PA, UNITED STATES

Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003171267 US 2001-833117	A1	20030911 20010412	(9)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 59 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 20 Drawing Page(s)

LINE COUNT: 13208

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 7 OF 9 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER: 2003:181414 USPATFULL
TITLE: Albumin fusion proteins

INVENTOR(S): Rosen, Craig A., Laytonsville, MD, UNITED STATES
Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003125247	A1	20030703	
APPLICATION INFO.:	US 2001-833041	A1	20010412	(9)

			NUMBER	DATE	
		- - -			
PRIORITY	INFORMATION:	US	2000-256931P	20001221	(60)
		US	2000-199384P	20000425	(60)
		US	2000-229358P	20000412	(60)
		TT.			

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 20 Drawing Page(s)

LINE COUNT: 15235

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 8 OF 9 USPATFULL on STN

Coated particles, methods of making and using TI

A particle coated with a nonlamellar material such as a nonlamellar AΒ crystalline material, a nonlamellar amorphous material, or a nonlamellar semi-crystalline material includes an internal matrix core having at least one a nanostructured liquid phase, or at least on nanostructured liquid crystalline phase or a combination of the two is used for the delivery of active agents such as pharmaceuticals, nutrients, pesticides, etc. The coated particle can be fabricated by a variety of different techniques where the exterior coating is a nonlamellar material such as a nonlamellar crystalline material, a nonlamellar amorphous material, or a nonlamellar semi-crystalline material

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:159130 USPATFULL

TITLE:

Coated particles, methods of making and using

INVENTOR(S):

Anderson, David M., Colonial Heights, VA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003108743	A1	20030612	
	US 6638621	B2	20031028	
APPLICATION INFO.:	US 2002-170237	A1	20020613	(1

RELATED APPLN. INFO.:

10) Continuation-in-part of Ser. No. US 2000-297997, filed

on 16 Aug 2000, GRANTED, Pat. No. US 6482517 DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

WHITHAM, CURTIS & CHRISTOFFERSON, P.C., 11491 SUNSET

HILLS ROAD, SUITE 340, RESTON, VA, 20190

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

107

NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT:

5538

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 9 OF 9 USPATFULL on STN L13

Multifunctional protease inhibitors and their use in treatment of TΙ disease

Fusion proteins of protease inhibitors are provided, in AB particular fusion proteins of alpha 1-antitrypsin (AAT) and a second protease inhibitor, such as secretory leukocyte protease inhibitor (SLPI) or tissue inhibitor of metalloproteases (TIMP). Polynucleotides encoding the fusion proteins, vectors comprising such polynucleotides, and host cells containing such vectors are also provided. Methods of making the fusion proteins of the invention are also provide, as well as methods of using the fusion proteins, for example to inhibit protease activity in a biological sample or in the treatment of an individual suffering from, or at risk for, a disease or disorder involving unwanted protease activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:106306 USPATFULL

TITLE:

Multifunctional protease inhibitors and their use in

treatment of disease

INVENTOR(S):

Barr, Philip J., Oakland, CA, UNITED STATES Gibson, Helen, Oakland, CA, UNITED STATES

Pemberton, Philip, San Francisco, CA, UNITED STATES

NUMBER KIND DATE PATENT INFORMATION: US 2003073217 A1 20030417 A1 20011218 (10)

NUMBER

US 2001-25514

APPLICATION INFO.:

DATE

PRIORITY INFORMATION: US 2000-256699P 20001218 (60) US 2001-331966P 20011120 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: MORRISON & FOERSTER LLP, 755 PAGE MILL RD, PALO ALTO,

CA, 94304-1018

NUMBER OF CLAIMS: 35 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 6 Drawing Page(s)

LINE COUNT: 3252

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

L8 8080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

L9 5 S L8 AND L1

L10 378 S L8 AND FUSION

L11 221 S L10 AND ALBUMIN

L12 5 S L11 AND ALBUMIN FRAGMENT

L13 9 S L11 AND SHELF-LIFE

=> s l11 and N-terminus fusion

L14 0 L11 AND N-TERMINUS FUSION

=> s l11 and C-terminus fusion

L15 0 L11 AND C-TERMINUS FUSION

=> d l11 ti abs ibib 1-25

L11 ANSWER 1 OF 221 USPATFULL on STN

TI Biospecific contrast agents

Methods and apparatuses for detecting a condition of a sample (including cervical cancers and pre-cancers) through reflectance and/or fluorescence imaging. A sample is obtained. One or more metallic nanoparticles and/or one or more quantum dots are obtained. The one or more metallic nanoparticles and/or one or more quantum dots are coupled to one or more biomarkers of the sample that are associated with the condition. A reflectance and/or fluorescence image of the sample is then taken. The image(s) exhibit characteristic optical scattering from the one or more metallic nanoparticles and/or characteristic fluorescence excitation from the one or more quantum dots to signal the presence of the one or more biomarkers. In this way, the condition can be readily screened or diagnosed.

ACCESSION NUMBER:

2004:31276 USPATFULL

TITLE:

Biospecific contrast agents

INVENTOR(S):

Sokolov, Konstantin, Austin, TX, UNITED STATES Korgel, Brian A., Round Rock, TX, UNITED STATES Ellington, Andrew D., Austin, TX, UNITED STATES Richards-Kortum, Rebecca, Austin, TX, UNITED STATES

NUMBER KIND DATE ______ US 2004023415 A1 20040205 US 2003-382136 A1 20030305 (10) PATENT INFORMATION: APPLICATION INFO.:

> NUMBER DATE _____

PRIORITY INFORMATION:

US 2002-361924P 20020305 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE: Michael C. Barrett, Esq., FULBRIGHT & JAWORSKI, L.L.P.,

600 Congress Avenue, Suite 2400, Austin, TX, 78701

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT: 3948

L11 ANSWER 2 OF 221 USPATFULL on STN

Biochips for characterizing biological processes TI

This invention includes biochips for analysis of a variety of molecules, AΒ cell components and cells. Embodiments of this invention include devices and methods for the parallel and/or nearly parallel processing of biological analytes. Biochips can comprise a substrate, Raman signal-enhancing structures, and receptors selective and/or specific for the analyte(s) to be assayed. Biochips can be read using a Raman reader and can provide for rapid, sensitive, direct assays for physiological and/or pathophysiological conditions of interest.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:31155 USPATFULL

TITLE: INVENTOR(S): Biochips for characterizing biological processes Kreimer, David I., Berkeley, CA, UNITED STATES Nufert, Thomas H., Walnut Creek, CA, UNITED STATES

Ginzburg, Lev, Fremont, CA, UNITED STATES Yevin, Oleg A., Oakland, CA, UNITED STATES

KIND DATE NUMBER -----

PATENT INFORMATION: APPLICATION INFO.:

US 2004023293 A1 20040205 US 2002-294385 A1 20021114 (10)

Continuation-in-part of Ser. No. US 2001-925189, filed RELATED APPLN. INFO.: on 8 Aug 2001, PENDING Continuation-in-part of Ser. No.

US 2001-815909, filed on 23 Mar 2001, PENDING

Continuation-in-part of Ser. No. US 2000-670453, filed

on 26 Sep 2000, PENDING

NUMBER DATE

PRIORITY INFORMATION:

US 1999-156195P 19990927 (60) US 2001-336445P 20011114 (60)

DOCUMENT TYPE: Utility FILE SEGMENT:

APPLICATION

Sheldon R. Meyer, FLIESLER DUBB MEYER & LOVEJOY LLP, LEGAL REPRESENTATIVE: Fourth Floor, Four Embarcadero Center, San Francisco,

CA, 94111-4156

NUMBER OF CLAIMS:

40

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

37 Drawing Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 3 OF 221 USPATFULL on STN

Proteases TI

The invention provides human proteases (PRTS) and polynucleotides which AB identify and encode PRTS. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with aberrant expression of PRTS.

ACCESSION NUMBER:

2004:31105 USPATFULL

TITLE:

INVENTOR(S):

Proteases Henry, Yue, Sunnyvale, CA, UNITED STATES

Elliott, Vicki S, San Jose, CA, UNITED STATES R Gandhi, Ameena, San Francisco, CA, UNITED STATES Lal, Preeti G, Santa Clara, CA, UNITED STATES Au-Young, Janice, Brisbane, CA, UNITED STATES Tribouley, Catherine M, San Francisco, CA, UNITED

STATES Delegeane, Angelo M, Milpitas, CA, UNITED STATES Baughn, Mariah R, San Leandro, CA, UNITED STATES Nguyen, Danniel B, San Jose, CA, UNITED STATES

Lee, Ernestine A, Albany, CA, UNITED STATES Hafalia, April J A, Daly City, CA, UNITED STATES Khan, Farrah A, Des Plaines, IL, UNITED STATES Chawla, Narinder K, Union City, CA, UNITED STATES Yao, Monique G, Carmel, IN, UNITED STATES

Lu, Dyung Aina M, San Jose, CA, UNITED STATES Arvizu, Chandra S, San Jose, CA, UNITED STATES Tang, Y Tom, San Jose, CA, UNITED STATES

Walsh, Roderick T, Canterbury, UNITED KINGDOM Azimzai, Yalda, Oakland, CA, UNITED STATES

Lu, Yan, Palo Alto, CA, UNITED STATES Ramkumar, Jayalaxmi, Fremont, CA, UNITED STATES Xu, Yuming, Mountain View, CA, UNITED STATES Reddy, Roopa, Sunnyvale, CA, UNITED STATES

Das, Debopriya, Mountain View, CA, UNITED STATES Kearney, Liam, San Francisco, CA, UNITED STATES Kallick, Deborah A, Galveston, TX, UNITED STATES

	NUMBER	KIND	DATE	
US	2004023243	A1	20040205	
US	2003-311035	A1	20030519	(10)
WO	2001-US19178		20010613	

DOCUMENT TYPE: FILE SEGMENT:

Utility

PATENT INFORMATION: APPLICATION INFO.:

APPLICATION

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

NUMBER OF CLAIMS: 116 EXEMPLARY CLAIM: LINE COUNT: 8891

L11 ANSWER 4 OF 221 USPATFULL on STN

Novel human gene relating to respiratory diseases, obesity, and ΤI inflammatory bowel disease

This invention relates to genes identified from human chromosome AΒ 20p13-p12, which are associated with various diseases, including asthma. The invention also relates to the nucleotide sequences of these genes, isolated nucleic acids comprising these nucleotide sequences, and isolated polypeptides or peptides encoded thereby. The invention further relates to vectors and host cells comprising the disclosed nucleotide sequences, or fragments thereof, as well as antibodies that bind to the encoded polypeptides or peptides. Also related are ligands that modulate

the activity of the disclosed genes or gene products. In addition, the invention relates to methods and compositions employing the disclosed nucleic acids, polypeptides or peptides, antibodies, and/or ligands for use in diagnostics and therapeutics for asthma and other diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:31077 USPATFULL

TITLE:

INVENTOR(S):

Novel human gene relating to respiratory diseases,

obesity, and inflammatory bowel disease

Keith, Tim, Bedford, MA, UNITED STATES

Little, Randall D., Newtonville, MA, UNITED STATES

Eerdewegh, Paul Van, Weston, MA, UNITED STATES

Dupuis, Josee, Newton, MA, UNITED STATES

Del Mastro, Richard G., Norfolk, MA, UNITED STATES

Simon, Jason, Westfield, NJ, UNITED STATES Allen, Kristina, Hopkinton, MA, UNITED STATES Pandit, Sunil, Gaithersburg, MD, UNITED STATES

KIND DATE NUMBER _____

PATENT INFORMATION: APPLICATION INFO.:

US 2004023215 A1 20040205 US 2002-126022 A1 20020419 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2001-834597, filed

on 13 Apr 2001, PENDING Continuation-in-part of Ser. No. US 2000-548797, filed on 13 Apr 2000, PENDING

NUMBER DATE -----------

PRIORITY INFORMATION: US 1999-129391P 19990413 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT: LEGAL REPRESENTATIVE:

APPLICATION MORGAN & FINNEGAN, L.L.P., 345 Park Avenue, New York,

NY, 10154-0053

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

73

NUMBER OF DRAWINGS:

157 Drawing Page(s)

LINE COUNT:

20001

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 5 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TΤ

The present invention relates to novel proteins. More specifically, AB isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

ACCESSION NUMBER:

2004:25127 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

NUMBER KIND DATE ______

PATENT INFORMATION:

US 2004018969

A1 20040129

APPLICATION INFO.:	US 2001-764875	A1 20010117 (9)	
	NUMBER	DATE	
PRIORITY INFORMATION:	US 2000-179065P US 2000-180628P US 2000-214886P US 2000-217487P US 2000-225758P US 2000-220963P	20000131 (60) 20000204 (60) 20000628 (60) 20000711 (60) 20000814 (60) 20000726 (60)	
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DOCUMENT TYPE:

FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

24 1

LINE COUNT:

38235

ANSWER 6 OF 221 USPATFULL on STN L11

TT Molecules for diagnostics and therapeutics

The present invention provides purified human polynucleotides for AΒ diagnostics and therapeutics (dithp). Also encompassed are the polypeptides (DITHP) encoded by dithp. The invention also provides for the use of dithp, or complements, oligonucleotides, or fragments thereof in diagnostic assays. The invention further provides for vectors and host cells containing dithp for the expression of DITHP. The invention additionally provides for the use of isolated and purified DITHP to induce antibodies and to screen libraries of compounds and the use of anti-DITHP antibodies in diagnostic assays. Also provided are microarrays containing dithp and methods of use.

ACCESSION NUMBER:

TITLE:

INVENTOR (S):

2004:18785 USPATFULL

Molecules for diagnostics and therapeutics Hodgson, David M., Ann Arbor, MI, UNITED STATES Lincoln, Stephen E., Potomac, MD, UNITED STATES Russo, Frank D., Sunnyvale, CA, UNITED STATES Albany, Peter A., Berkeley, CA, UNITED STATES Banville, Steve C., Sunnyvale, CA, UNITED STATES Bratcher, Shawn R., Mountain View, CA, UNITED STATES Dufour, Gerard E., Castro Valley, CA, UNITED STATES Cohen, Howard J., Palo Alto, CA, UNITED STATES Rosen, Bruce H., Menlo Park, CA, UNITED STATES Chalup, Michael S., Livingston, TX, UNITED STATES Jackson, Jennifer L., Santa Cruz, CA, UNITED STATES Jones, Anissa L., San Jose, CA, UNITED STATES Yu, Jimmy Y., Fremont, CA, UNITED STATES Greenawalt, Lila B., San Jose, CA, UNITED STATES Panzer, Scott R., Sunnyvale, CA, UNITED STATES Roseberry Lincoln, Ann M., Potomac, MD, UNITED STATES Wright, Rachel J., Merivale, NEW ZEALAND

PATENT ASSIGNEE(S):

PATENT

Daniels, Susan E., Mountain View, CA, UNITED STATES Incyte Corporation, Palo Alto, CA, UNITED STATES (U.S.

corporation)

	NUMBER	KIND	DATE	
INFORMATION:	US 2004014087	A1	20040122	
ATION INFO.:	US 2003-378029	A1	20030228	

APPLICA (10) . RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 2001-980285, filed on 30 Nov 2001, PENDING A 371 of International Ser. No. WO 2000-US15404, filed on 31 May 2000, PENDING

DATE NUMBER ______ US 1999-147500P 19990805 (60) PRIORITY INFORMATION: 19990805 (60) US 1999-147542P 19990805 (60) US 1999-147541P 19990805 (60) US 1999-147824P 19990805 (60) US 1999-147547P 19990805 (60) US 1999-147530P 19990805 (60) US 1999-147536P 19990805 (60) US 1999-147520P 19990805 (60) US 1999-147527P 19990805 (60) US 1999-147549P 19990804 (60) US 1999-147377P 19990804 (60) US 1999-147436P US 1999-137411P 19990603 (60) 19990603 (60) US 1999-137396P US 1999-137417P 19990603 (60) 19990603 (60) US 1999-137337P 19990602 (60) US 1999-137173P 19990602 (60) US 1999-137114P 19990602 (60) US 1999-137259P 19990602 (60) US 1999-137113P 19990602 (60) US 1999-137260P 19990602 (60) US 1999-137258P 19990602 (60) US 1999-137109P 19990601 (60) US 1999-137161P Utility

DOCUMENT TYPE: .

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 14819

L11 ANSWER 7 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TI

The present invention relates to novel proteins. More specifically, AB isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

ACCESSION NUMBER:

2004:18737 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

PATENT ASSIGNEE(S):

Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES, 20850 (U.S. corporation)

	NUMBER	KIND	DATE.	
N:	US 2004014039	A1	20040122	
_	TTC 2002 150057	Δ1	20020531	

PATENT INFORMATION APPLICATION INFO.:

(10) US 2002-158057

RELATED APPLN. INFO.:

Continuation of Ser. No. US 2001-764890, filed on 17

Jan 2001, PENDING

	NUMBER	DATE
PRIORITY INFORMATION:	US 2000-179065P US 2000-180628P US 2000-214886P US 2000-217487P US 2000-225758P US 2000-225758P US 2000-217496P US 2000-218290P US 2000-218290P US 2000-225757P US 2000-226868P US 2000-216647P US 2000-216880P US 2000-216880P US 2000-251869P US 2000-2353834P US 2000-2353834P US 2000-2353834P US 2000-235834P US 2000-236369P US 2000-234274P US 2000-224518P US 2000-224519P US 2000-2245186P US 2000-2244617P US 2000-225268P US 2000-2251868P US 2000-229344P	20000131 (60) 20000204 (60) 20000628 (60) 20000711 (60) 20000814 (60) 20000714 (60) 20000814 (60) 20000814 (60) 20000814 (60) 20000814 (60) 20000814 (60) 20000707 (60) 20000814 (60) 20000707 (60) 20000814 (60) 20000927 (60) 20000921 (60) 20000921 (60) 20000814 (60) 20000921 (60) 20000814 (60) 20000921 (60) 20000814 (60) 20000921 (60) 20000814 (60) 20000921 (60) 20000814 (60) 20000929 (60) 20000814 (60) 20000929 (60) 20001101 (60) 20001208 (60) 20001020 (60) 20001101 (60) 20000929 (60) 20001208 (60) 20001208 (60) 20001208 (60) 20001208 (60) 20001208 (60) 20001208 (60) 20001208 (60) 20000925 (60) 20000925 (60) 20000925 (60) 20000925 (60) 20000925 (60) 20000925 (60) 20000925 (60)
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DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

Utility

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

26776 LINE COUNT:

L11 ANSWER 8 OF 221 USPATFULL on STN

Albumin fusion proteins TI

The present invention encompasses albumin fusion AB proteins. Nucleic acid molecules encoding the albumin

fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using

these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating,

preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL Albumin fusion proteins

TITLE: INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

DATE NUMBER KIND _____ US 2004010134 20040115

PATENT INFORMATION: APPLICATION INFO.:

A1 A1 20010412 (9) US 2001-833245

NUMBER DATE _____ US 2000-256931P 20001221 (60) PRIORITY INFORMATION: 20000425 (60) US 2000-199384P US 2000-229358P 20000412 (60)

Utility DOCUMENT TYPE: FILE SEGMENT: APPLICATION

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 9 OF 221 USPATFULL on STN

7 Human ovarian and ovarian cancer associated proteins ΤI

This invention relates to newly identified ovarian or ovarian cancer AB related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "ovarian cancer antigens", and the use of such ovarian antigens for detecting disorders of the reproductive system, particularly the presence of ovarian cancer and ovarian cancer metastases. This invention relates to ovarian cancer antigens as well as vectors, host cells, antibodies directed to ovarian cancer antigens and the recombinant methods and synthetic methods for producing the same. Also provided are diagnostic methods for detecting, treating, preventing and/or prognosing disorders related to the ovary, including ovarian cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of ovarian cancer antigens of the invention. The present invention further relates to inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

INVENTOR(S):

2004:13598 USPATFULL

TI

AΒ

7 Human ovarian and ovarian cancer associated proteins Birse, Charles E., North Potomac, MD, UNITED STATES

20010316

Rosen, Craig A., Laytonsville, MD, UNITED STATES

NUMBER KIND DATE US 2004010121 A1 US 2003-333900 A1 20040115 20030124 (10) WO 2001-US8585

PATENT INFORMATION: APPLICATION INFO.:

DOCUMENT TYPE:

Utility APPLICATION

FILE SEGMENT: LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

23 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 16023 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 10 OF 221 USPATFULL on STN

Use of bioactive glass compositions to stimulate osteoblast production Compositions comprising bioactive glass compositions or extracts thereof which include ions in an appropriate concentration and ratio that they enhance osteoblast production, and methods of preparation and use thereof, are disclosed. The compositions can be included in implantable devices that are capable of inducing tissue formation in autogeneic, allogeneic and xenogeneic implants, for example as coatings and/or matrix materials. Examples of such devices include prosthetic implants, sutures, stents, screws, plates, tubes, and the like. Aqueous extracts of the bioactive glass compositions, which extracts are capable of stimulating osteoblast production, are also disclosed. The compositions can be used, for example, to induce local tissue formation from a progenitor cell in a mammal, for accelerating allograft repair in a mammal, for promoting in vivo integration of an implantable prosthetic device to enhance the bond strength between the prosthesis and the existing target tissue at the joining site, and for treating tissue degenerative conditions.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13078 USPATFULL

TITLE:

Use of bioactive glass compositions to stimulate

osteoblast production

INVENTOR(S):

Hench, Larry L, London, UNITED KINGDOM Polak, Julia M, London, UNITED KINGDOM Buttery, Lee D.k., London, UNITED KINGDOM

Xynos, Ioannis D, Nafplion, GREECE

Maroothynaden, Jason, London, UNITED KINGDOM

KIND DATE NUMBER _____

20040115 A1 PATENT INFORMATION: US 2004009598

A1 20030707 (10) US 2003-332731 APPLICATION INFO .:

20010711 WO 2001-US21801

Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT:

BURNS DOANE SWECKER & MATHIS L L P, POST OFFICE BOX LEGAL REPRESENTATIVE:

1404, ALEXANDRIA, VA, 22313-1404

NUMBER OF CLAIMS: 1 EXEMPLARY CLAIM: LINE COUNT: 1301

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 11 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI

The present invention relates to novel polynucleotides associated with AB the plasma membrane, the polypeptides encoded by these polynucleotides herein collectively referred to as "plasma membrane associated antigens," and antibodies that immunospecifically bind these polypeptides, and the use of such plasma membrane associated polynucleotides, antigens, and antibodies for detecting, treating, preventing and/or prognosing disorders related to these novel polypeptides. More specifically, isolated nucleic acid molecules are provided encoding novel plasma membrane associated polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing these plasma membrane associated polynucleotides, polypeptides, and/or antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the novel polypeptides of the invention. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The invention further relates to methods and/or compositions for inhibiting or promoting the production and/or function of the polypeptides of the invention.

ACCESSION NUMBER:

2004:12971 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Birse, Charles E., North Potomac, MD, UNITED STATES

Rosen, Craig A., Laytonsville, MD, UNITED STATES

KIND DATE NUMBER _____

PATENT INFORMATION: APPLICATION INFO .:

US 2004009491 A1 20040115 US 2002-264237 A1 20021004 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. WO 2001-US16450, filed

on 18 May 2001, PENDING

DATE NUMBER _____

PRIORITY INFORMATION:

US 2000-205515P 20000519 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24 EXEMPLARY CLAIM: 1 LINE COUNT: 18144

L11 ANSWER 12 OF 221 USPATFULL on STN

ΤI Nucleic acids, proteins, and antibodies

The present invention relates to novel musculoskeletal system related AB polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "musculoskeletal system antigens," and the use of such musculoskeletal system antigens for detecting disorders of

the musculoskeletal system, particularly the presence of cancer and cancer metastases. More specifically, isolated musculoskeletal system associated nucleic acid molecules are provided encoding novel musculoskeletal system associated polypeptides. Novel musculoskeletal system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human musculoskeletal system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the musculoskeletal system, including cancer of musculoskeletal tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:12968 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES, 20850 (U.S. corporation)

NUMBER KIND DATE ______

PATENT INFORMATION: APPLICATION INFO.:

PATENT ASSIGNEE(S):

US 2004009488 A1 20040115 US 2002-242515 A1 20020913 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-764877, filed on 17

Jan 2001, PENDING

NUMBER DATE

PRIORITY INFORMATION:

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US 2001-259678P
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Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 24

LINE COUNT: 32038

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 13 OF 221 USPATFULL on STN

TI Methods for the treatment of carcinoma

The invention concerns compositions and methods for the diagnosis and treatment of neoplastic cell growth and proliferation in mammals, including humans. The invention is based upon the identification of genes that are amplified in the genome of tumor cells, such as renal cell carcinoma. Such gene amplification is expected to be associated with the overexpression of the gene product as compared to normal cells of the same tissue type and contribute to tumorigenesis. Accordingly, the proteins encoded by the amplified genes are believed to be useful targets for the diagnosis and/or treatment (including prevention) of certain cancers, such as renal cell carcinoma, and may act as predictors

of the prognosis of tumor treatment. The present invention is directed to novel methods of diagnosing and treating tumor, such as renal cell carcinoma or Wilms tumor.

ACCESSION NUMBER:

2004:12653 USPATFULL

TITLE:

Methods for the treatment of carcinoma

INVENTOR(S):

Gerritsen, Mary E., San Mateo, CA, UNITED STATES

Peale, Franklin V., JR., San Carlos, CA, UNITED STATES

Wu, Thomas D., San Francisco, CA, UNITED STATES

PATENT ASSIGNEE(S):

GENENTECH, INC. (U.S. corporation)

NUMBER KIND ______ A1 20040115 A1 20030221 US 2004009171 PATENT INFORMATION:

APPLICATION INFO.:

US 2003-372683 20030221 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2002-271690, filed

on 16 Oct 2002, PENDING

NUMBER DATE

PRIORITY INFORMATION:

US 2001-344534P 20011018 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

GENENTECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA,

94080

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT:

AΒ

57 1 6662

L11 ANSWER 14 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI

The present invention relates to novel ovarian related polynucleotides, the polypeptides encoded by these polynucleotides herein collectively referred to as "ovarian antigens," and antibodies that immunospecifically bind these polypeptides, and the use of such ovarian polynucleotides, antigens, and antibodies for detecting, treating, preventing and/or prognosing disorders of the reproductive system, particularly disorders of the ovaries and/or breast, including, but not limited to, the presence of ovarian and/or breast cancer and ovarian and/or breast cancer metastases. More specifically, isolated ovarian nucleic acid molecules are provided encoding novel ovarian polypeptides. Novel ovarian polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human ovarian polynucleotides, polypeptides, and/or antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the ovaries and/or breast, including ovarian and/or breast cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The invention further relates to methods and/or compositions for inhibiting or promoting the production and/or function of the polypeptides of the invention.

ACCESSION NUMBER:

2004:7345 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Birse, Charles E., North Potomac, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES

NUMBER	KIND	DATE	
US 2004005579	A1	20040108	
US 2002-264049	A1	20021004	(

APPLICATION INFO .: RELATED APPLN. INFO.:

PATENT INFORMATION:

Continuation-in-part of Ser. No. WO 2001-US18569, filed

on 7 Jun 2001, PENDING

DATE NUMBER _____ US 2000-209467P 20000607 (60)

PRIORITY INFORMATION: DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

18130 LINE COUNT:

L11 ANSWER 15 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI

The present invention relates to novel proteins. More specifically, AB isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

PATENT ASSIGNEE(S):

2004:7343 USPATFULL

Nucleic acids, proteins, and antibodies

INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES (U.S. corporation)

NUMBER KIND DATE US 2004005577 A1 20040108 US 2002-242747 A1 20020913 (10)

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

Continuation of Ser. No. US 2001-764881, filed on 17

20000814 (60)

20001208 (60)

20000927 (60)

Jan 2001, PENDING

			NUMBER	DATE	
PRIORITY	INFORMATION:	US	2000-179065P	20000131	(60)
		US	2000-180628P	20000204	(60)
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US 2000-205515P
US 2001-259678P
                    20010105 (60)
Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24 EXEMPLARY CLAIM: 1 LINE COUNT: 27694

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 16 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies

ΤI AB

The present invention relates to novel cardiovascular system related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "cardiovascular system antigens," and the use of such cardiovascular system antigens for detecting disorders of the cardiovascular system, particularly the presence of cancer of cardiovascular system tissues and cancer metastases. More specifically, isolated cardiovascular system associated nucleic acid molecules are provided encoding novel cardiovascular system associated polypeptides. Novel cardiovascular system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human cardiovascular system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the cardiovascular system, including cancer of cardiovascular system tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:7341 USPATFULL

NUMBER

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD, UNITED

DATE

PATENT ASSIGNEE(S):

STATES, 20850 (U.S. corporation) KIND

PATENT INFORMATION: APPLICATION INFO.:

_____ -----US 2004005575 A1 20040108

RELATED APPLN. INFO.:

A1 20020826 US 2002-227577 (10)

Continuation of Ser. No. US 2002-91504, filed on 7 Mar 2002, PENDING Continuation of Ser. No. US 2001-764869,

20000921 (60)

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filed on 17 Jan 2001, ABANDONED

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US 2001-259678P
Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24
EXEMPLARY CLAIM: 1
LINE COUNT: 28742

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 17 OF 221 USPATFULL on STN

TI Functional MRI agents for cancer imaging

AB The invention relates to novel magnetic resonance imaging contrast agents for imaging cancer.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

2004:4285 USPATFULL ACCESSION NUMBER:

Functional MRI agents for cancer imaging TITLE:

Meade, Thomas J., Altadena, CA, United States INVENTOR(S): Fraser, Scott, La Canada, CA, United States

Jacobs, Russell, Arcadia, CA, United States

Research Corporation Technologies, Inc., Tucson, AZ, PATENT ASSIGNEE(S):

United States (U.S. corporation)

KIND DATE NUMBER _____

PATENT INFORMATION:

B1 20040106 US 6673333

US 2000-715859 APPLICATION INFO.:

20001117 (9)

DATE NUMBER ______

US 2000-201816P 20000504 (60) PRIORITY INFORMATION:

Utility DOCUMENT TYPE: GRANTED FILE SEGMENT:

Hartley, Michael G. PRIMARY EXAMINER:

Dorsey & Whitney LLP, Silva, Robin M., Kossiak, Renee LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM:

7 Drawing Figure(s); 5 Drawing Page(s) NUMBER OF DRAWINGS:

2422 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 18 OF 221 USPATFULL on STN

50 human secreted proteins ΤI

The present invention relates to novel human secreted proteins and AB isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:2568 USPATFULL

TITLE:

50 human secreted proteins

INVENTOR(S):

Moore, Paul A., Germantown, MD, UNITED STATES Ruben, Steven M., Olney, MD, UNITED STATES LaFleur, David W., Washington, DC, UNITED STATES Shi, Yanggu, Gaithersburg, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES Olsen, Henrik S., Gaithersburg, MD, UNITED STATES Ebner, Reinhard, Gaithersburg, MD, UNITED STATES Brewer, Laurie A., St. Paul, MN, UNITED STATES Human Genome Sciences, Inc., Rockville, MD (U.S.

PATENT ASSIGNEE(S):

corporation)

KIND DATE NUMBER ______

PATENT INFORMATION: APPLICATION INFO .:

US 2004002591 A1 20040101 US 2002-47021 A1 20020117 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2000-722329, filed on 28 Nov 2000, PENDING Continuation of Ser. No. US

1999-262109, filed on 4 Mar 1999, ABANDONED

Continuation-in-part of Ser. No. WO 1998-US18360, filed

on 3 Sep 1998, PENDING

PRIORITY INFORMATION:

US 2001-262066P 20010118 (60) 19970905 (60) US 1997-57626P

19970905 (60) US 1997-57663P

US 1997-57669P 19970905 (60)

19970912 (60) US 1997-58666P 19970912 (60) US 1997-58667P

19970912 (60) US 1997-58973P

US 1997-58974P 19970912 (60) 19980622 (60) US 1998-90112P

Utility DOCUMENT TYPE:

FILE SEGMENT: APPLICATION

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 23 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT: 33379

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 19 OF 221 USPATFULL on STN

ΤI Novel human gene relating to respiratory diseases, obesity, and

inflammatory bowel disease

This invention relates to genes identified from human chromosome AB20p13-p12, which are associated with various diseases, including asthma. The invention also relates to the nucleotide sequences of these genes, isolated nucleic acids comprising these nucleotide sequences, and isolated polypeptides or peptides encoded thereby. The invention further relates to vectors and host cells comprising the disclosed nucleotide sequences, or fragments thereof, as well as antibodies that bind to the encoded polypeptides or peptides. Also related are ligands that modulate the activity of the disclosed genes or gene products. In addition, the invention relates to methods and compositions employing the disclosed nucleic acids, polypeptides or peptides, antibodies, and/or ligands for use in diagnostics and therapeutics for asthma and other diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:2447 USPATFULL

TITLE:

Novel human gene relating to respiratory diseases,

obesity, and inflammatory bowel disease

INVENTOR(S):

Keith, Tim, Bedford, MA, UNITED STATES

Little, Randall D., Newtonville, MA, UNITED STATES

Eerdewegh, Paul Van, Weston, MA, UNITED STATES

Dupuis, Josee, Newton, MA, UNITED STATES

Del Mastro, Richard G., Norfolk, MA, UNITED STATES Simon, Jason, Westfield, NJ, UNITED STATES

Allen, Kristin, Hopkinton, MA, UNITED STATES Pandit, Sunil, Gaithersburg, MD, UNITED STATES

NUMBER	KIND	DATE

PATENT INFORMATION:

US 2004002470 A1 20040101

APPLICATION INFO.:

A1 20021017 (10) US 2002-277216

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2002-126022, filed on 19 Apr 2002, PENDING Continuation-in-part of Ser. No. US 2001-834597, filed on 13 Apr 2001, PENDING Continuation-in-part of Ser. No. US 2000-548797, filed

on 13 Apr 2000, PENDING

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

MORGAN & FINNEGAN, L.L.P., 345 PARK AVENUE, NEW YORK,

NY, 10154

NUMBER OF CLAIMS:

45

EXEMPLARY CLAIM:

1

162 Drawing Page(s) NUMBER OF DRAWINGS:

15810 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 20 OF 221 USPATFULL on STN

Detection and modulation of Slit and roundabount (Robo) mediated TI

angiogenesis and uses thereof

This invention is generally in the field of methods for diagnosis, AB treatment and prevention of various disorders involving the Slit2 mediated angiogenesis.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:335332 USPATFULL

TITLE:

Detection and modulation of Slit and roundabount (Robo)

mediated angiogenesis and uses thereof

INVENTOR (S):

Geng, Jian-Guo, Portage, MI, UNITED STATES

DATE NUMBER KIND

PATENT INFORMATION: APPLICATION INFO.:

US 2003236210 A1 20031225 US 2003-386386 A1 20030310

A1 20030310 (10)

NUMBER DATE -----

PRIORITY INFORMATION:

US 2002-362485P 20020308 (60)

DOCUMENT TYPE:

Utility

APPLICATION

FILE SEGMENT: LEGAL REPRESENTATIVE:

Peng Chen, Morrison & Foerster LLP, Suite 500, 3811

Valley Centre Drive, San Diego, CA, 92130-2332

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

4 Drawing Page(s)

NUMBER OF DRAWINGS: LINE COUNT:

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1337

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 21 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TI

The present invention relates to novel excretory system related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "excretory system antigens," and the use of such excretory system antigens for detecting disorders of the excretory system, particularly the presence of cancer of excretory system tissues and cancer metastases. More specifically, isolated excretory system associated nucleic acid molecules are provided encoding novel excretory system associated polypeptides. Novel excretory system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human excretory system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the excretory system, including cancer of excretory system tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

2003:334955 USPATFULL ACCESSION NUMBER:

TITLE:

Nucleic acids, proteins, and antibodies

Rosen, Craig A., Laytonsville, MD, UNITED STATES INVENTOR (S):

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

PATENT ASSIGNEE(S):

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

Human Genome Sciences, Inc., Rockville, MD, 20850 (U.S. corporation)

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Continuation of S	Ser. No	. US 2001-764897,	filed on 17

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DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT:

22457

24

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 22 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TI

The present invention relates to novel proteins. More specifically, AΒ isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:334953 USPATFULL

TITLE: INVENTOR (S): Nucleic acids, proteins, and antibodies Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES Birse, Charles E., North Potomac, MD, UNITED STATES

PATENT ASSIGNEE(S):

Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES (U.S. corporation)

NUMBER KIND DATE ------ -----US 2003235829 A1 20031225 PATENT INFORMATION: 20020826 APPLICATION INFO .: US 2002-227646 **A1**

RELATED APPLN. INFO.:

Continuation of Ser. No. US 2001-860670, filed on 21 May 2001, PENDING Continuation-in-part of Ser. No. WO

2001-US1346, filed on 17 Jan 2001, PENDING

NUMBER

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US 2000-205515P
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Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24 EXEMPLARY CLAIM: 1

LINE COUNT: 20415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 23 OF 221 USPATFULL on STN

TI Compositions and methods for systemic inhibition of cartilage degradation

AB Methods and compositions for inhibiting articular cartilage degradation. The compositions preferably include multiple chondroprotective agents, including at least one agent that promotes cartilage anabolic activity and at least one agent that inhibits cartilage catabolism. The compositions may also include one or more pain and inflammation

inhibitory agents. The compositions may be administered systemically, such as to treat patients at risk of cartilage degradation at multiple joints, and suitably may be formulated in a carrier or delivery vehicle that is targeted to the joints. Alternatively the compositions may be injected or infused directly into the joint.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:334713 USPATFULL

TITLE:

Compositions and methods for systemic inhibition of

cartilage degradation

INVENTOR (S):

Demopulos, Gregory A., Mercer Island, WA, UNITED STATES Palmer, Pamela Pierce, San Francisco, CA, UNITED STATES

Herz, Jeffrey M., Mill Creek, WA, UNITED STATES

PATENT ASSIGNEE(S):

Omeros Corporation (U.S. corporation)

NUMBER	KIND	DATE

PATENT INFORMATION:

US 2003235589 A1 20031225

APPLICATION INFO.: RELATED APPLN. INFO.:

US 2003-356649 A1 20030131 (10)

Continuation-in-part of Ser. No. US 2002-31546, filed on 18 Jan 2002, PENDING A 371 of International Ser. No.

WO 2000-US19864, filed on 21 Jul 2000, PENDING

Continuation-in-part of Ser. No. US 2001-839633, filed on 20 Apr 2001, PENDING Continuation-in-part of Ser. No. WO 1999-US26330, filed on 5 Nov 1999, PENDING Continuation-in-part of Ser. No. WO 1999-US24625, filed

on 20 Oct 1999, PENDING

		NUMBER	DATE
PRIORITY	INFORMATION:	US 2002-353552P	20020201

US 2002-353552P 20020201 (60) US 1999-144904P 19990721 (60) US 1998-107256P 19981105 (60) US 1998-105026P 19981020 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

OMEROS MEDICAL SYSTEMS, INC., 1420 FIFTH AVENUE, SUITE

2675, SEATTLE, WA, 98101

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 155

NUMBER OF DRAWINGS:

9 Drawing Page(s)

LINE COUNT:

6575

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L11 ANSWER 24 OF 221 USPATFULL on STN
- TI Nucleic acids, proteins, and antibodies

The present invention relates to novel endocrine related polynucleotides AΒ and the polypeptides encoded by these polynucleotides herein collectively known as "endocrine antigens," and the use of such endocrine antigens for detecting disorders of the endocrine system, particularly the presence of cancers of the endocrine system and endocrine cancer metastases. More specifically, isolated endocrine associated nucleic acid molecules are provided encoding novel endocrine associated polypeptides. Novel endocrine polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human endocrine associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the endocrine system, including cancers of the endocrine system, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the

production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:330759 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Ruben, Steven M., Olney, MD, UNITED STATES Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD (U.S.

corporation)

NUMBER KIND DATE _____

PATENT INFORMATION:

PATENT ASSIGNEE(S):

US 2003232975 A1 20031218

APPLICATION INFO.:

20020214 US 2002-74024 A1 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-764895, filed on 17

Jan 2001, ABANDONED

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Utility
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DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

24

EXEMPLARY CLAIM:

21828

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 25 OF 221 USPATFULL on STN

TI Proteases

The invention provides human proteases (PRTS) and polynucleotides which AB identify and encode PRTS. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with aberrant expression of PRTS.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:330138 USPATFULL

TITLE:

Proteases

INVENTOR(S):

Delegeane, Angelo M., Milpitas, CA, UNITED STATES Gandhi, Ameena R., San Francisco, CA, UNITED STATES Hafalia, April J. A., Santa Clara, CA, UNITED STATES Lu, Dyung Aina M., San Jose, CA, UNITED STATES Arvizu, Chandra S., San Jose, CA, UNITED STATES Tribouley, Catherine M., San Francisco, CA, UNITED STATES

Das, Debopriya, Mountain View, CA, UNITED STATES Kallick, Deborah A., Portola Valley, CA, UNITED STATES Nguyen, Danniel B., San Jose, CA, UNITED STATES Lee, Ernestine A., Castro Valley, CA, UNITED STATES Khan, Farrah A., Glen View, IL, UNITED STATES Yue, Henry, Sunnyvale, CA, UNITED STATES Au-Young, Janice, Brisbane, CA, UNITED STATES Griffin, Jennifer A., Fremont, CA, UNITED STATES Policky, Jennifer L., San Jose, CA, UNITED STATES

Ramkumar, Jayalaxmi, Fremont, CA, UNITED STATES Yang, Junming, San Jose, CA, UNITED STATES Thangavelu, Kavitha, Mountain View, CA, UNITED STATES Ding, Li, Creve Coeur, MO, UNITED STATES Kearney, Liam, San Francisco, CA, UNITED STATES Baughn, Mariah R., San Leandro, CA, UNITED STATES Borowsky, Mark L., Redwood City, CA, UNITED STATES Sanjanwala, Madhusudan, Los Altos, CA, UNITED STATES Yao, Monique G., Carmel, IN, UNITED STATES Burford, Neil, Durham, CT, UNITED STATES Chawla, Narinder K., Union City, CA, UNITED STATES Lal, Preeti G., Santa Clara, CA, UNITED STATES Lee, Sally, San Jose, CA, UNITED STATES Todd, Stephen, San Francisco, CA, UNITED STATES Lo, Terence P., Foster City, CA, UNITED STATES Tang, Y. Tom, San Jose, CA, UNITED STATES Elliott, Vicki S., San Jose, CA, UNITED STATES Azimzai, Yalda, Oakland, CA, UNITED STATES Lu, Yan, Palo Alto, CA, UNITED STATES

PATENT ASSIGNEE(S):

Incyte Genomics, Inc., Palo Alto, CA (U.S. corporation)

	NUMBER	KIND	DATE			
PATENT INFORMATION:	US 2003232349	A1	20031218			
APPLICATION INFO .:	US 2002-274639	A1	20021018	(10)		
RELATED APPLN. INFO.:	Continuation of	Ser. No	. WO 2001-	US22397,	filed	on 17
	Jul 2001, PENDIN	1G				

			NUMBER	DATE	
PRIORITY	INFORMATION:		2000-220063P 2000-221680P	20000721 20000728	(60) (60)
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		US	2000-224717P	20000811	(60)
		US	2000-225988P	20000816	(60)
		US	2000-227568P	20000823	(60)
DOCUMENT	TYPE:	Uti	lity		

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

LINE COUNT:

8959

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

NEWS PHONE

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CAS World Wide Web Site (general information)

* STN Columbus

FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004

=> file medline, uspatful, dgene, embase, wpids COST IN U.S. DOLLARS

SINCE FILE TOTAL SESSION ENTRY

FULL ESTIMATED COST

0.21 0.21

FILE 'MEDLINE' ENTERED AT 13:52:47 ON 06 FEB 2004

FILE 'USPATFULL' ENTERED AT 13:52:47 ON 06 FEB 2004 CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE 'WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004 COPYRIGHT (C) 2004 THOMSON DERWENT

=> s albumin fusion proteins

2835 ALBUMIN FUSION PROTEINS

=> s cerebus protein

1 CEREBUS PROTEIN

=> s 11 and 12

0 L1 AND L2

=> s (cerebus protein) and albumin

O (CEREBUS PROTEIN) AND ALBUMIN

=> s 12 and fusion

0 L2 AND FUSION

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ANSWER 1 OF 1 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L2

Human and murine cerebus-like proteins - used for treating tissue defects TΙ and degenerative nerve conditions.

ΔN 1999-106054 [09] WPIDS

CR 2003-298696 [29]

9901553 A UPAB: 20030505 AΒ

A novel isolated DNA sequence comprises a DNA sequence selected from: (a) nucleotides beginning at # 1, 52, 55, 58, 61, 64, 67, 70, 73, 121, 256, 259, 262, 265, 268, 171, or 484 and ending at # 723 or 801 of the 804 bp DNA sequence given in the specification; and (b) sequences which hybridise to (a) under stringent hybridisation conditions and encode a protein which exhibits cerebus activity. Also claimed are: (1) an isolated DNA sequence comprising nucleotides encoding amino acids beginning at #1, 18 to 25, 41, 85 to 91 or 152, and ending at #241 or 267 of the 267 amino acid sequence given in the specification; (2) a vector comprising either of the above DNA molecules in operative association with an expression control sequence; (3) an isolated DNA molecule comprising nucleotides 268-801 of the 272 amino acid sequence given in the specification (sic), or naturally occurring allelic sequences of it; (4) a vector comprising the DNA of (4) in operative association with an expression control sequence; (5) an isolated DNA molecule encoding mammalian cerebus protein , comprising nucleotides 268-801 of the 804 bp DNA sequence given in the

specification; (6) a vector comprising the DNA of (5) in operative association with an expression control sequence; (7) a host cell

transformed with the vector of (2), (4) or (6); (8) a purified mammalian cerebus protein comprising the 267 amino acid sequence given in the specification; (9) a purified mammalian cerebus protein comprising residues 90-267 of the 272 amino acid sequence given in the specification; and (10) antibodies to the cerebus protein of (8) or (9).

USE - The host cell of (7) can be used to produce the mammalian cerebus proteins (claimed). Compositions containing the protein can be used in the formation of neurons and related neural cells and tissues, such as Schwann cells, glial cells, and astrocytes, as well as liver, pancreas, lung, heart, kidney, spleen, stomach, and cardiac tissue and cells. They may also be used to treat precursor or stem cells. The compositions can also be used for treating tissue defects, and healing and maintenance of various types of tissues and wounds. The mammalian cerebus protein containing compositions may also be used to treat or prevent degenerate nerve conditions such as Parkinson's disease, Alzheimer's disease, and Lou Gehrig's disease. They can also be used to treat osteoporosis, rheumatoid arthritis, osteoarthritis, and other abnormalities of connective tissue, or of other organs or tissues. Dwq.0/0

ACCESSION NUMBER:

1999-106054 [09] WPIDS

CROSS REFERENCE:

2003-298696 [29]

DOC. NO. CPI:

C1999-031758

TITLE:

Human and murine cerebus-like proteins - used for treating tissue defects and degenerative nerve

conditions.

DERWENT CLASS:

B04 D16

INVENTOR(S):

DEROBERTIS, E M; FOLLETTIE, M

PATENT ASSIGNEE(S):

(GEMY) GENETICS INST INC; (REGC) UNIV CALIFORNIA

COUNTRY COUNT:

PATENT INFORMATION:

PATENT 1	NO.	KIND	DATE	WEEK	LA	PG

WO 9901553 A1 19990114 (199909) * EN 50

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG UZ VN YU ZW

57

A 19990125 (199923) AU 9878140

A 19990810 (199938) US 5935852

A1 20000628 (200035) EN EP 1012278

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

MX 2000000242 A1 20010601 (200235)

JP 2002511762 W 20020416 (200242)

B 20020620 (200252) AU 749031

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 9901553	A1	WO 1998-US11462	19980603
AU 9878140	A	AU 1998-78140	19980603
US 5935852	A	US 1997-887997	19970703
EP 1012278	A1	EP 1998-926263	19980603
		WO 1998-US11462	19980603
MX 2000000242	A1	MX 2000-242	20000105
JP 2002511762	W	WO 1998-US11462	19980603
		JP 1999-507147	19980603
AU 749031	В	AU 1998-78140	19980603

FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 9878140 EP 1012278 JP 2002511762 AU 749031	A Based on A1 Based on W Based on B Previous Publ. Based on	WO 9901553 WO 9901553 WO 9901553 . AU 9878140 WO 9901553

PRIORITY APPLN. INFO: US 1997-887997 19970703

=> file biosis

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 20.32 20.53

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004 COPYRIGHT (C) 2004 BIOLOGICAL ABSTRACTS INC.(R)

FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 4 February 2004 (20040204/ED)

FILE RELOADED: 19 October 2003.

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

=> s 12

0 CEREBUS

1361492 PROTEIN

L6 0 CE

O CEREBUS PROTEIN (CEREBUS (W) PROTEIN)

=> file medline, uspatful, dgene, embase, wpids, biosis, japio, fsta, jicst COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION 0.85 21.38

FULL ESTIMATED COST

FILE 'MEDLINE' ENTERED AT 14:00:26 ON 06 FEB 2004

FILE 'USPATFULL' ENTERED AT 14:00:26 ON 06 FEB 2004 CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE 'JAPIO' ENTERED AT 14:00:26 ON 06 FEB 2004 COPYRIGHT (C) 2004 Japanese Patent Office (JPO) - JAPIO

FILE 'FSTA' ENTERED AT 14:00:26 ON 06 FEB 2004 COPYRIGHT (C) 2004 International Food Information Service

FILE 'JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004 COPYRIGHT (C) 2004 Japan Science and Technology Agency (JST)

=> s 12

1 L2 L7

=> d 17 ti abs ibib tot

ANSWER 1 OF 1 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L7

Human and murine cerebus-like proteins - used for treating tissue defects TI and degenerative nerve conditions.

MΔ 1999-106054 [09] WPIDS

CR 2003-298696 [29]

9901553 A UPAB: 20030505 AB

A novel isolated DNA sequence comprises a DNA sequence selected from: (a) nucleotides beginning at # 1, 52, 55, 58, 61, 64, 67, 70, 73, 121, 256, 259, 262, 265, 268, 171, or 484 and ending at # 723 or 801 of the 804 bp DNA sequence given in the specification; and (b) sequences which hybridise to (a) under stringent hybridisation conditions and encode a protein which exhibits cerebus activity. Also claimed are: (1) an isolated DNA sequence comprising nucleotides encoding amino acids beginning at #1, 18 to 25, 41, 85 to 91 or 152, and ending at #241 or 267 of the 267 amino acid sequence given in the specification; (2) a vector comprising either of the above DNA molecules in operative association with an expression control sequence; (3) an isolated DNA molecule comprising nucleotides 268-801 of the 272 amino acid sequence given in the specification (sic), or naturally occurring allelic sequences of it; (4) a vector comprising the DNA of (4) in operative association with an expression control sequence; (5) an isolated DNA molecule encoding mammalian cerebus protein , comprising nucleotides 268-801 of the 804 bp DNA sequence given in the specification; (6) a vector comprising the DNA of (5) in operative association with an expression control sequence; (7) a host cell transformed with the vector of (2), (4) or (6); (8) a purified mammalian cerebus protein comprising the 267 amino acid sequence given in the specification; (9) a purified mammalian cerebus protein comprising residues 90-267 of the 272 amino acid sequence given in the specification; and (10) antibodies to the cerebus protein of (8) or (9).

USE - The host cell of (7) can be used to produce the mammalian cerebus proteins (claimed). Compositions containing the protein can be used in the formation of neurons and related neural cells and tissues, such as Schwann cells, glial cells, and astrocytes, as well as liver, pancreas, lung, heart, kidney, spleen, stomach, and cardiac tissue and cells. They may also be used to treat precursor or stem cells. The compositions can also be used for treating tissue defects, and healing and maintenance of various types of tissues and wounds. The mammalian cerebus protein containing compositions may also be used to treat or prevent degenerate nerve conditions such as Parkinson's disease, Alzheimer's disease, and Lou Gehrig's disease. They can also be used to treat osteoporosis, rheumatoid arthritis, osteoarthritis, and other abnormalities of connective tissue, or of other organs or tissues.

Dwq.0/0

ACCESSION NUMBER:

1999-106054 [09] WPIDS

CROSS REFERENCE:

2003-298696 [29]

DOC. NO. CPI:

C1999-031758

TITLE:

Human and murine cerebus-like proteins - used for treating tissue defects and degenerative nerve

conditions.

DERWENT CLASS:

B04 D16

INVENTOR(S):

DEROBERTIS, E M; FOLLETTIE, M

PATENT ASSIGNEE(S):

(GEMY) GENETICS INST INC; (REGC) UNIV CALIFORNIA

COUNTRY COUNT: 83

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9901553 A1 19990114 (199909)* EN 50

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL

OA PT SD SE SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG

MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG

57

UZ VN YU ZW

AU 9878140 A 19990125 (199923)

US 5935852 A 19990810 (199938)

EP 1012278 A1 20000628 (200035) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

MX 2000000242 A1 20010601 (200235)

JP 2002511762 W 20020416 (200242)

AU 749031 B 20020620 (200252)

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 9901553	A1		1998-US11462	19980603
AU 9878140	A		1998-78140	19980603
US 5935852	A	US	1997-887997	19970703
EP 1012278	A1	ΕP	1998-926263	19980603
		WO	1998-US11462	19980603
MX 2000000242	A1	MX	2000-242	20000105
JP 2002511762	W	WO	1998-US11462	19980603
		JP	1999-507147	19980603
AU 749031	В	ΑU	1998-78140	19980603

FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 9878140 EP 1012278 JP 2002511762 AU 749031	A Based on Al Based on W Based on B Previous Publ. Based on	WO 9901553 WO 9901553 WO 9901553 AU 9878140 WO 9901553

PRIORITY APPLN. INFO: US 1997-887997 19970703

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

=> s TIMP-1 or tissue inhibitor metalloproteinase-1

5 FILES SEARCHED...

L8 8080 TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

=> s 18 and 11

5 L8 AND L1

=> d 19 ti abs ibib tot

L9 ANSWER 1 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL Albumin fusion proteins

INVENTOR(S):

TITLE:

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

NUMBER	KIND	DATE	
 JS 2004010134 JS 2001-833245	A1 A1	20040115 20010412	(9)

DOCUMENT TYPE: FILE SEGMENT: Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 29 1

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 2 OF 5 USPATFULL on STN

TI Albumin fusion proteins

AB The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic

acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003219875	A1	20031127	(9)
APPLICATION INFO.:	US 2001-833118	A1	20010412	

NUMBER DATE

PRIORITY INFORMATION: US 2000-256931P 20001221 (60)

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT:

15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 3 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL Albumin fusion proteins

INVENTOR (S):

TTTLE:

Ballance, David J., Berwyn, PA, UNITED STATES Sleep, Darrell, West Bridgford, UNITED KINGDOM Prior, Christopher P., Rosemont, PA, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003199043	A1	20031023	
APPLICATION INFO.:	US 2001-832501	A1	20010412	(9)

NUMBER DATE

PRIORITY INFORMATION:

US 2000-256931P

20001221 (60)

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

60

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 4 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:244853 USPATFULL Albumin fusion proteins

TITLE: INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

		NUMBER	KIND	DATE	•
PATENT INFORMATION:	• •	2003171267 2001-833117	A1 A1	20030911	(9)
APPLICATION INFO.:	US	2001-83311/	ΑI	20010412	(9)

		NUMBER	DATE
PRIORITY I	NFORMATION:	US 2000-199384P 20	001221 (60) 000425 (60) 000412 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

LEGAL REPRESENTATIVE:

APPLICATION

RO

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

59

NUMBER OF DRAWINGS:

20 Drawing Page(s)

LINE COUNT:

13208

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L9 ANSWER 5 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion

proteins of the invention.

Albumin fusion proteins

The present invention encompasses albumin fusion

TI

AB

CAS INDEXING IS AVAILABLE FOR THIS PATENT. 2003:181414 USPATFULL ACCESSION NUMBER: Albumin fusion proteins TITLE: Rosen, Craig A., Laytonsville, MD, UNITED STATES INVENTOR(S): Haseltine, William A., Washington, DC, UNITED STATES KIND DATE NUMBER US 2003125247 A1 20030703 US 2001-833041 A1 20010412 PATENT INFORMATION: APPLICATION INFO.: DATE NUMBER _____ US 2000-256931P 20001221 (60) PRIORITY INFORMATION: US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60) Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE: ROCKVILLE, MD, 20850 NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 20 Drawing Page(s) NUMBER OF DRAWINGS: 15235 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. => d his (FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004) FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004 2835 S ALBUMIN FUSION PROTEINS L11 S CEREBUS PROTEIN L2L3 0 S L1 AND L2 0 S (CEREBUS PROTEIN) AND ALBUMIN L40 S L2 AND FUSION L5 FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004 L6 0 S L2 FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004 L7 1 S L2 8080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1 L8L95 S L8 AND L1 => s 18 and fusion 378 L8 AND FUSION L10 => s 110 and albumin 221 L10 AND ALBUMIN => s l11 and albumin fragment 5 L11 AND ALBUMIN FRAGMENT => d l12 ti abs ibib tot L12 ANSWER 1 OF 5 USPATFULL on STN

proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2004010134 US 2001-833245	A1 A1	20040115 20010412	(9)

NUMBER DATE

PRIORITY INFORMATION:

US 2000-256931P 20001221 (60)

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

1

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 2 OF 5 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003219875	A1	20031127	
APPLICATION INFO.:	US 2001-833118	A1	20010412	(9)

NUMBER DATE

US 2000-256931P 20001221 (60) PRIORITY INFORMATION:

US 2000-199384P 20000425 (60) 20000412 (60) US 2000-229358P

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 3 OF 5 USPATFULL on STN

Albumin fusion proteins TТ

The present invention encompasses albumin fusion AΒ proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL Albumin fusion proteins

TITLE: INVENTOR (S):

Ballance, David J., Berwyn, PA, UNITED STATES Sleep, Darrell, West Bridgford, UNITED KINGDOM Prior, Christopher P., Rosemont, PA, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003199043	A1	20031023	
APPLICATION INFO.:	US 2001-832501	A1	20010412	(9)

NUMBER DATE US 2000-256931P 20001221 (60) PRIORITY INFORMATION: US 2000-199384P 20000425 (60) 20000412 (60) US 2000-229358P

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 60 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT:

14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 4 OF 5 USPATFULL on STN L12

Albumin fusion proteins ΤI

The present invention encompasses albumin fusion AB proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using

these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:244853 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003171267 US 2001-833117	A1 A1	20030911	(9)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 59 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 20 Drawing Page(s)

LINE COUNT: 13208

13208

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 5 OF 5 USPATFULL on STN

TI Albumin fusion proteins
AB The present invention en

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:181414 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

•	NUMBER	KIND DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003125247 US 2001-833041	A1 20030703 A1 20010412	(9)
	NUMBER	DATE	
PRIORITY INFORMATION:	US 2000-256931P US 2000-199384P US 2000-229358P	20001221 (60) 20000425 (60) 20000412 (60)	

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

29

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

20 Drawing Page(s)

LINE COUNT:

15235

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d his

(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

L8 8080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

L9 5 S L8 AND L1

L10 378 S L8 AND FUSION

L11 . 221 S L10 AND ALBUMIN

L12 5 S L11 AND ALBUMIN FRAGMENT

=> s l11 and shelf-life

L13 9 L11 AND SHELF-LIFE

=> d l13 ti abs ibib tot

L13 ANSWER 1 OF 9 USPATFULL on STN

TI Biospecific contrast agents

Methods and apparatuses for detecting a condition of a sample (including cervical cancers and pre-cancers) through reflectance and/or fluorescence imaging. A sample is obtained. One or more metallic nanoparticles and/or one or more quantum dots are obtained. The one or more metallic nanoparticles and/or one or more quantum dots are coupled to one or more biomarkers of the sample that are associated with the condition. A reflectance and/or fluorescence image of the sample is then taken. The image(s) exhibit characteristic optical scattering from the one or more metallic nanoparticles and/or characteristic fluorescence excitation from the one or more quantum dots to signal the presence of the one or more biomarkers. In this way, the condition can be readily screened or diagnosed.

ACCESSION NUMBER:

2004:31276 USPATFULL

TITLE:

Biospecific contrast agents

INVENTOR (S):

Sokolov, Konstantin, Austin, TX, UNITED STATES Korgel, Brian A., Round Rock, TX, UNITED STATES Ellington, Andrew D., Austin, TX, UNITED STATES Richards-Kortum, Rebecca, Austin, TX, UNITED STATES

NUMBER KIND DATE

PATENT INFORMATION:

US 2004023415

20040205

APPLICATION INFO.:

US 2003-382136

20030305 (10)

NUMBER

DATE ______

A1

A1

PRIORITY INFORMATION:

US 2002-361924P

20020305 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

Michael C. Barrett, Esq., FULBRIGHT & JAWORSKI, L.L.P.,

600 Congress Avenue, Suite 2400, Austin, TX, 78701

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT:

3948

L13 ANSWER 2 OF 9 USPATFULL on STN

Albumin fusion proteins TI

AB

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

		NUMBER	KIND	DATE	
ATENT INFORMATION:	ບຣ	2004010134	A1	20040115	
PPLICATION INFO.:	US	2001-833245	A1	20010412	(9)

NUMBER DATE ______ US 2000-256931P 20001221 (60) PRIORITY INFORMATION: US 2000-199384P 20000425 (60) US 2000-199384P US 2000-229358P 20000412 (60)

Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

25066 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 3 OF 9 USPATFULL on STN

Nanoporous particle with a retained target TI

Porous nanostructured materials, such as porous nanostructured liquid AB and liquid crystalline particles or materials, incorporate a target substantially within the material which selectively binds a chemical of interest which can diffusion within the porous nanostructured material and be bound by the target. The porous nanostructured materials can be dispersed as particles in a medium in which said chemical of interest is located with low turbidity. Markers which detect binding of said

chemical of interest can be maintained in the medium separate and apart from the target, and any active compound (e.g., an enzyme) associated therewith by the porous nanostructured material, such that detectable changes in the marker only result when the active compounds diffuse out of the porous nanostructured materials after the chemical of interest binds to the target.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER: 2003:330129 USPATFULL

TITLE:

Nanoporous particle with a retained target

INVENTOR(S):

Anderson, David, Colonial Heights, VA, UNITED STATES

KIND DATE NUMBER ______ US 2003232340 A1 20031218

PATENT INFORMATION:

APPLICATION INFO.:

US 2002-170214

A1 20020613 (10)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

WHITHAM, CURTIS & CHRISTOFFERSON, P.C., 11491 SUNSET

HILLS ROAD, SUITE 340, RESTON, VA, 20190

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

2 Drawing Page(s)

LINE COUNT:

2555

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 4 OF 9 USPATFULL on STN

Albumin fusion proteins ΤI

AΒ

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating,

preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:312278 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003219875	A1	20031127	(9)
APPLICATION INFO.:	US 2001-833118	A1	20010412	

NUMBER DATE

US 2000-256931P 20001221 (60) PRIORITY INFORMATION:

US 2000-199384P 20000425 (60) US 2000-229358P 20000412 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

29

EXEMPLARY CLAIM:

1

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

15415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 5 OF 9 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:282700 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Ballance, David J., Berwyn, PA, UNITED STATES Sleep, Darrell, West Bridgford, UNITED KINGDOM Prior, Christopher P., Rosemont, PA, UNITED STATES Sadeghi, Homayoun, Doylestown, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003199043 US 2001-832501	A1 A1	20031023 20010412	(9)
	NUMBER	DA'	TE	
PRIORITY INFORMATION:	US 2000-256931P US 2000-199384P US 2000-229358P	2000 2000 2000	1 1	
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DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

60 1

NUMBER OF DRAWINGS:

18 Drawing Page(s)

LINE COUNT:

14339

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L13 ANSWER 6 OF 9 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:244853 USPATFULL

TITLE:

Albumin fusion proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Sadeghi, Homayoun, Doylestown, PA, UNITED STATES

Prior, Christopher P., Rosemont, PA, UNITED STATES Turner, Andrew J., Eagleville, PA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003171267 US 2001-833117			(9)
	NUMBER	DA'	ΓE	
PRIORITY INFORMATION:	US 2000-256931P US 2000-199384P US 2000-229358P	2000	0425 (60)	
DOCUMENT TYPE: FILE SEGMENT: LEGAL REPRESENTATIVE:	Utility APPLICATION HUMAN GENOME SCIE ROCKVILLE, MD, 20		NC, 9410 K	EY WEST AVENUE,
NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS: LINE COUNT: CAS INDEXING IS AVAILAB	13208			

L13 ANSWER 7 OF 9 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:181414 USPATFULL

TITLE: INVENTOR(S):

AB

Albumin fusion proteins Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 2003125247 US 2001-833041	A1 A1	20030703 20010412	
	NUMBER	DA	TE	
PRIORITY INFORMATION:	US 2000-256931P US 2000-199384P US 2000-229358P	2000	0425 (60)	
DOCUMENT TYPE: FILE SEGMENT: LEGAL REPRESENTATIVE:	Utility APPLICATION HUMAN GENOME SCIE ROCKVILLE, MD, 20		NC, 9410	KEY WEST AVENUE,
NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS: LINE COUNT: CAS INDEXING IS AVAILAB	29 1 20 Drawing Page(s 15235	1)		

L13 ANSWER 8 OF 9 USPATFULL on STN

Coated particles, methods of making and using ΤI

A particle coated with a nonlamellar material such as a nonlamellar AΒ crystalline material, a nonlamellar amorphous material, or a nonlamellar semi-crystalline material includes an internal matrix core having at least one a nanostructured liquid phase, or at least on nanostructured liquid crystalline phase or a combination of the two is used for the delivery of active agents such as pharmaceuticals, nutrients, pesticides, etc. The coated particle can be fabricated by a variety of different techniques where the exterior coating is a nonlamellar material such as a nonlamellar crystalline material, a nonlamellar amorphous material, or a nonlamellar semi-crystalline material

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:159130 USPATFULL

TITLE:

Coated particles, methods of making and using

INVENTOR (S):

Anderson, David M., Colonial Heights, VA, UNITED STATES

	NUMBER	KIND	DATE		
PATENT INFORMATION:	US 2003108743	A1	20030612		
	US 6638621	B2	20031028		
APPLICATION INFO.:	US 2002-170237	A1	20020613	(10)	
RELATED APPLN. INFO.:	Continuation-in-	part of	Ser. No.	US 2000-297997,	filed
•	on 16 Aug 2000,	GRANTED	, Pat. No.	. US 6482517	
DOCUMENT TYPE:	Utility				
FILE SEGMENT:	APPLICATION				

LEGAL REPRESENTATIVE:

WHITHAM, CURTIS & CHRISTOFFERSON, P.C., 11491 SUNSET

HILLS ROAD, SUITE 340, RESTON, VA, 20190

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

107

NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 9 OF 9 USPATFULL on STN

TΙ Multifunctional protease inhibitors and their use in treatment of

Fusion proteins of protease inhibitors are provided, in AB particular fusion proteins of alpha 1-antitrypsin (AAT) and a second protease inhibitor, such as secretory leukocyte protease inhibitor (SLPI) or tissue inhibitor of metalloproteases (TIMP). Polynucleotides encoding the fusion proteins, vectors comprising such polynucleotides, and host cells containing such vectors are also provided. Methods of making the fusion proteins of the invention are also provide, as well as methods of using the fusion proteins, for example to inhibit protease activity in a biological sample or in the treatment of an individual suffering from, or at risk for, a disease or disorder involving unwanted protease activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:106306 USPATFULL

TITLE:

Multifunctional protease inhibitors and their use in

treatment of disease

INVENTOR(S):

Barr, Philip J., Oakland, CA, UNITED STATES Gibson, Helen, Oakland, CA, UNITED STATES

Pemberton, Philip, San Francisco, CA, UNITED STATES

NUMBER KIND DATE ______ PATENT INFORMATION: US 2003073217 A1 20030417 APPLICATION INFO.: US 2001-25514 A1 20011218 (10)

> NUMBER DATE

PRIORITY INFORMATION: US 2000-256699P 20001218 (60) US 2001-331966P 20011120 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: MORRISON & FOERSTER LLP, 755 PAGE MILL RD, PALO ALTO,

CA, 94304-1018

NUMBER OF CLAIMS: 35 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 6 Drawing Page(s)

LINE COUNT: 3252

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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(FILE 'HOME' ENTERED AT 13:52:01 ON 06 FEB 2004)

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS' ENTERED AT 13:52:47 ON 06 FEB 2004

L1 2835 S ALBUMIN FUSION PROTEINS

L2 1 S CEREBUS PROTEIN

L3 0 S L1 AND L2

L4 0 S (CEREBUS PROTEIN) AND ALBUMIN

L5 0 S L2 AND FUSION

FILE 'BIOSIS' ENTERED AT 13:59:37 ON 06 FEB 2004

L6 0 S L2

FILE 'MEDLINE, USPATFULL, DGENE, EMBASE, WPIDS, BIOSIS, JAPIO, FSTA, JICST-EPLUS' ENTERED AT 14:00:26 ON 06 FEB 2004

L7 1 S L2

L8 8080 S TIMP-1 OR TISSUE INHIBITOR METALLOPROTEINASE-1

L9 5 S L8 AND L1

L10 378 S L8 AND FUSION

L11 221 S L10 AND ALBUMIN

L12 5 S L11 AND ALBUMIN FRAGMENT

L13 9 S L11 AND SHELF-LIFE

=> s lll and N-terminus fusion

L14 0 L11 AND N-TERMINUS FUSION

=> s l11 and C-terminus fusion

L15 0 L11 AND C-TERMINUS FUSION

=> d l11 ti abs ibib 1-25

L11 ANSWER 1 OF 221 USPATFULL on STN

TI Biospecific contrast agents

Methods and apparatuses for detecting a condition of a sample (including cervical cancers and pre-cancers) through reflectance and/or fluorescence imaging. A sample is obtained. One or more metallic nanoparticles and/or one or more quantum dots are obtained. The one or more metallic nanoparticles and/or one or more quantum dots are coupled to one or more biomarkers of the sample that are associated with the condition. A reflectance and/or fluorescence image of the sample is then taken. The image(s) exhibit characteristic optical scattering from the one or more metallic nanoparticles and/or characteristic fluorescence excitation from the one or more quantum dots to signal the presence of the one or more biomarkers. In this way, the condition can be readily screened or diagnosed.

ACCESSION NUMBER:

2004:31276 USPATFULL

TITLE:

Biospecific contrast agents

INVENTOR(S):

Sokolov, Konstantin, Austin, TX, UNITED STATES Korgel, Brian A., Round Rock, TX, UNITED STATES Ellington, Andrew D., Austin, TX, UNITED STATES Richards-Kortum, Rebecca, Austin, TX, UNITED STATES

KIND DATE NUMBER _____ US 2004023415 A1 US 2003-382136 A1 20040205 20030305 (10)

DATE NUMBER ______

PRIORITY INFORMATION:

PATENT INFORMATION:

APPLICATION INFO .:

Utility

US 2002-361924P 20020305 (60)

DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

Michael C. Barrett, Esq., FULBRIGHT & JAWORSKI, L.L.P.,

600 Congress Avenue, Suite 2400, Austin, TX, 78701

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

11 Drawing Page(s)

LINE COUNT:

3948

L11 ANSWER 2 OF 221 USPATFULL on STN

Biochips for characterizing biological processes TI

This invention includes biochips for analysis of a variety of molecules, AB cell components and cells. Embodiments of this invention include devices and methods for the parallel and/or nearly parallel processing of biological analytes. Biochips can comprise a substrate, Raman signal-enhancing structures, and receptors selective and/or specific for the analyte(s) to be assayed. Biochips can be read using a Raman reader and can provide for rapid, sensitive, direct assays for physiological and/or pathophysiological conditions of interest.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:31155 USPATFULL

TITLE: INVENTOR(S): Biochips for characterizing biological processes Kreimer, David I., Berkeley, CA, UNITED STATES Nufert, Thomas H., Walnut Creek, CA, UNITED STATES

Ginzburg, Lev, Fremont, CA, UNITED STATES Yevin, Oleg A., Oakland, CA, UNITED STATES

NUMBER KIND DATE -----

PATENT INFORMATION: APPLICATION INFO .:

US 2004023293 A1 20040205 US 2002-294385 A1 20021114 (10)

Continuation-in-part of Ser. No. US 2001-925189, filed RELATED APPLN. INFO.: on 8 Aug 2001, PENDING Continuation-in-part of Ser. No.

US 2001-815909, filed on 23 Mar 2001, PENDING

Continuation-in-part of Ser. No. US 2000-670453, filed

on 26 Sep 2000, PENDING

DATE NUMBER _____

PRIORITY INFORMATION:

US 1999-156195P 19990927 (60) US 2001-336445P 20011114 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

Sheldon R. Meyer, FLIESLER DUBB MEYER & LOVEJOY LLP, Fourth Floor, Four Embarcadero Center, San Francisco,

CA. 94111-4156

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

40 1

NUMBER OF DRAWINGS:

37 Drawing Page(s)

LINE COUNT:

3572

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 3 OF 221 USPATFULL on STN

Proteases TI

The invention provides human proteases (PRTS) and polynucleotides which AB identify and encode PRTS. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with aberrant expression of PRTS.

ACCESSION NUMBER:

2004:31105 USPATFULL

TITLE:

INVENTOR(S):

Proteases Henry, Yue, Sunnyvale, CA, UNITED STATES

Elliott, Vicki S, San Jose, CA, UNITED STATES R Gandhi, Ameena, San Francisco, CA, UNITED STATES Lal, Preeti G, Santa Clara, CA, UNITED STATES Au-Young, Janice, Brisbane, CA, UNITED STATES Tribouley, Catherine M, San Francisco, CA, UNITED

STATES Delegeane, Angelo M, Milpitas, CA, UNITED STATES Baughn, Mariah R, San Leandro, CA, UNITED STATES Nguyen, Danniel B, San Jose, CA, UNITED STATES Lee, Ernestine A, Albany, CA, UNITED STATES Hafalia, April J A, Daly City, CA, UNITED STATES Khan, Farrah A, Des Plaines, IL, UNITED STATES Chawla, Narinder K, Union City, CA, UNITED STATES Yao, Monique G, Carmel, IN, UNITED STATES Lu, Dyung Aina M, San Jose, CA, UNITED STATES Arvizu, Chandra S, San Jose, CA, UNITED STATES Tang, Y Tom, San Jose, CA, UNITED STATES Walsh, Roderick T, Canterbury, UNITED KINGDOM Azimzai, Yalda, Oakland, CA, UNITED STATES Lu, Yan, Palo Alto, CA, UNITED STATES Ramkumar, Jayalaxmi, Fremont, CA, UNITED STATES Xu, Yuming, Mountain View, CA, UNITED STATES Reddy, Roopa, Sunnyvale, CA, UNITED STATES Das, Debopriya, Mountain View, CA, UNITED STATES Kearney, Liam, San Francisco, CA, UNITED STATES Kallick, Deborah A, Galveston, TX, UNITED STATES

	NUMBER	KIND	DATE
US	2004023243	A1	20040205
US	2003-311035	A1	20030519

PATENT INFORMATION: APPLICATION INFO .:

Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

20010613

(10)

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

116 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 8891 LINE COUNT:

L11 ANSWER 4 OF 221 USPATFULL on STN

Novel human gene relating to respiratory diseases, obesity, and ΤI

WO 2001-US19178

inflammatory bowel disease

This invention relates to genes identified from human chromosome AΒ 20p13-p12, which are associated with various diseases, including asthma. The invention also relates to the nucleotide sequences of these genes, isolated nucleic acids comprising these nucleotide sequences, and isolated polypeptides or peptides encoded thereby. The invention further relates to vectors and host cells comprising the disclosed nucleotide sequences, or fragments thereof, as well as antibodies that bind to the encoded polypeptides or peptides. Also related are ligands that modulate the activity of the disclosed genes or gene products. In addition, the invention relates to methods and compositions employing the disclosed nucleic acids, polypeptides or peptides, antibodies, and/or ligands for use in diagnostics and therapeutics for asthma and other diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

INVENTOR(S):

2004:31077 USPATFULL

TITLE:

Novel human gene relating to respiratory diseases,

obesity, and inflammatory bowel disease

Keith, Tim, Bedford, MA, UNITED STATES

Little, Randall D., Newtonville, MA, UNITED STATES

Eerdewegh, Paul Van, Weston, MA, UNITED STATES

Dupuis, Josee, Newton, MA, UNITED STATES

Del Mastro, Richard G., Norfolk, MA, UNITED STATES

Simon, Jason, Westfield, NJ, UNITED STATES Allen, Kristina, Hopkinton, MA, UNITED STATES Pandit, Sunil, Gaithersburg, MD, UNITED STATES

NUMBER KIND DATE -----

PATENT INFORMATION: APPLICATION INFO.:

US 2004023215 A1 20040205 US 2002-126022 A1 20020419 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2001-834597, filed on 13 Apr 2001, PENDING Continuation-in-part of Ser. No. US 2000-548797, filed on 13 Apr 2000, PENDING

DATE NUMBER

PRIORITY INFORMATION:

Utility.

US 1999-129391P 19990413 (60)

DOCUMENT TYPE:

APPLICATION

FILE SEGMENT: LEGAL REPRESENTATIVE:

MORGAN & FINNEGAN, L.L.P., 345 Park Avenue, New York,

NY, 10154-0053

NUMBER OF CLAIMS:

73

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

157 Drawing Page(s)

LINE COUNT:

20001

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 5 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI

The present invention relates to novel proteins. More specifically, AB isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

ACCESSION NUMBER:

2004:25127 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

NUMBER KIND DATE US 2004018969 A1 20040129

PATENT INFORMATION:

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APPLICATION INFO.:	US 2001-764875	A1 20010117	(9)
APPLICATION INFO			(2)
	NUMBER	DATE ·	
PRIORITY INFORMATION:	US 2000-179065P	20000131 (60)	
	US 2000-180628P	20000204 (60)	
	US 2000-214886P	20000628 (60)	
	US 2000-217487P US 2000-225758P	20000711 (60) 20000814 (60)	
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	US 2000-217496P	20000711 (60)	
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	US 2000-225757P	20000814 (60)	
•	US 2000-226868P US 2000-216647P	20000822 (60) 20000707 (60)	
	US 2000-225267P	20000814 (60)	
	US 2000-216880P	20000707 (60)	
	US 2000-225270P	20000814 (60)	
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	US 2000-235834P US 2000-234274P	20000927 (60)	
	US 2000-234223P	20000921 (60)	
	US 2000-228924P	20000830 (60)	
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	US 2000-241809P	20001020 (60)	
	US 2000-249299P	20001117 (60)	
	US 2000-236327P	20000929 (60)	
	US 2000-241785P	20001020 (60) 20001101 (60)	
	US 2000-244617P US 2000-225268P	20001101 (60)	
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	US 2000-251856P	20001208 (60)	
	US 2000-251868P	20001208 (60)	
	US 2000-229344P US 2000-234997P	20000901 (60) 20000925 (60)	
	US 2000-234337F	20000923 (60)	
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	US 2000-229513P	20000905 (60)	
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•	US 2000-236367P	20000929 (60)	
	US 2000-237039P	20001002 (60)	
	US 2000-237038P	20001002 (60)	
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	US 2000-237037P	20001002 (60)	•
	US 2000-237040P	20001002 (60)	
	US 2000-240960P	20001020 (60)	
	US 2000-239935P	20001013 (60)	
	US 2000-239937P US 2000-241787P	20001013 (60) 20001020 (60)	
	US 2000-241787P	20001020 (80)	
	US 2000-246532P	20001108 (60)	
	US 2000-249216P	20001117 (60)	
	US 2000-249210P	20001117 (60)	·
	US 2000-226681P US 2000-225759P	20000822 (60) 20000814 (60)	
	US 2000-225213P	20000814 (60)	
	US 2000-227182P	20000822 (60)	

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US 2000-225214P
                    20000814 (60)
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                    20000927 (60)
                    20000906 (60)
US 2000-230438P
US 2000-215135P
                    20000630 (60)
                    20000814 (60)
US 2000-225266P
                    20001117 (60)
US 2000-249218P
US 2000-249208P
                    20001117 (60)
                    20001117 (60)
US 2000-249213P
US 2000-249212P
                    20001117 (60)
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DOCUMENT TYPE:

Utility APPLICATION

FILE SEGMENT: LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

24 38235

LINE COUNT:

L11 ANSWER 6 OF 221 USPATFULL on STN

Molecules for diagnostics and therapeutics ΤI

The present invention provides purified human polynucleotides for AB diagnostics and therapeutics (dithp). Also encompassed are the polypeptides (DITHP) encoded by dithp. The invention also provides for the use of dithp, or complements, oligonucleotides, or fragments thereof in diagnostic assays. The invention further provides for vectors and host cells containing dithp for the expression of DITHP. The invention additionally provides for the use of isolated and purified DITHP to induce antibodies and to screen libraries of compounds and the use of anti-DITHP antibodies in diagnostic assays. Also provided are microarrays containing dithp and methods of use.

ACCESSION NUMBER:

TITLE:

INVENTOR (S):

2004:18785 USPATFULL

Molecules for diagnostics and therapeutics Hodgson, David M., Ann Arbor, MI, UNITED STATES Lincoln, Stephen E., Potomac, MD, UNITED STATES Russo, Frank D., Sunnyvale, CA, UNITED STATES Albany, Peter A., Berkeley, CA, UNITED STATES Banville, Steve C., Sunnyvale, CA, UNITED STATES Bratcher, Shawn R., Mountain View, CA, UNITED STATES Dufour, Gerard E., Castro Valley, CA, UNITED STATES Cohen, Howard J., Palo Alto, CA, UNITED STATES Rosen, Bruce H., Menlo Park, CA, UNITED STATES Chalup, Michael S., Livingston, TX, UNITED STATES Jackson, Jennifer L., Santa Cruz, CA, UNITED STATES Jones, Anissa L., San Jose, CA, UNITED STATES Yu, Jimmy Y., Fremont, CA, UNITED STATES Greenawalt, Lila B., San Jose, CA, UNITED STATES

Panzer, Scott R., Sunnyvale, CA, UNITED STATES Roseberry Lincoln, Ann M., Potomac, MD, UNITED STATES

Wright, Rachel J., Merivale, NEW ZEALAND

Daniels, Susan E., Mountain View, CA, UNITED STATES Incyte Corporation, Palo Alto, CA, UNITED STATES (U.S.

corporation)

PATENT ASSIGNEE(S):

NUMBER	KIND	DATE
US 2004014087	A1	20040122

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

A1 20030228 (10) US 2003-378029

Continuation-in-part of Ser. No. US 2001-980285, filed on 30 Nov 2001, PENDING A 371 of International Ser. No. WO 2000-US15404, filed on 31 May 2000, PENDING

DATE _____ 19990805 (60) US 1999-147500P PRIORITY INFORMATION: 19990805 (60) US 1999-147542P 19990805 (60) US 1999-147541P US 1999-147824P 19990805 (60) 19990805 (60) US 1999-147547P 19990805 (60) US 1999-147530P 19990805 (60) US 1999-147536P US 1999-147520P 19990805 (60) US 1999-147527P 19990805 (60) US 1999-147549P 19990805 (60) 19990804 (60) US 1999-147377P US 1999-147436P 19990804 (60) 19990603 (60) US 1999-137411P 19990603 (60) US 1999-137396P US 1999-137417P 19990603 (60) 19990603 (60) US 1999-137337P 19990602 (60) US 1999-137173P 19990602 (60) US 1999-137114P 19990602 (60) US 1999-137259P US 1999-137113P 19990602 (60) US 1999-137260P 19990602 (60) 19990602 (60) US 1999-137258P 19990602 (60) US 1999-137109P 19990601 (60) US 1999-137161P Utility

NUMBER

DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

NUMBER OF CLAIMS: 19 EXEMPLARY CLAIM: LINE COUNT: 14819

L11 ANSWER 7 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TI

The present invention relates to novel proteins. More specifically, AB isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

ACCESSION NUMBER:

2004:18737 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES, 20850 (U.S. corporation)

KIND NUMBER DATE A1 20040122 US 2004014039

PATENT INFORMATION: APPLICATION INFO.:

PATENT ASSIGNEE(S):

A1 20020531 US 2002-158057 (10)

RELATED APPLN. INFO.:

Continuation of Ser. No. US 2001-764890, filed on 17

Jan 2001, PENDING

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DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT:

1 26776

L11 ANSWER 8 OF 221 USPATFULL on STN

TI Albumin fusion proteins

The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disordrs or conditions using albumin fusion proteins of the invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13611 USPATFULL

TITLE:
INVENTOR(S):

Albumin fusion proteins Rosen, Craig A., Laytonsville, MD, UNITED STATES

Haseltine, William A., Washington, DC, UNITED STATES

NUMBER KIND DATE

PATENT INFORMATION: US 2004010134 A1 20040115

APPLICATION INFO.: US 2001-833245 A1 20010412 (9)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 25066

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 9 OF 221 USPATFULL on STN

TI 7 Human ovarian and ovarian cancer associated proteins

This invention relates to newly identified ovarian or ovarian cancer related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "ovarian cancer antigens",

and the use of such ovarian antigens for detecting disorders of the reproductive system, particularly the presence of ovarian cancer and ovarian cancer metastases. This invention relates to ovarian cancer antigens as well as vectors, host cells, antibodies directed to ovarian cancer antigens and the recombinant methods and synthetic methods for producing the same. Also provided are diagnostic methods for detecting, treating, preventing and/or prognosing disorders related to the ovary, including ovarian cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of ovarian cancer antigens of the invention. The present invention further relates to inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

INVENTOR(S):

2004:13598 USPATFULL

TITLE:

TI

AΒ

7 Human ovarian and ovarian cancer associated proteins Birse, Charles E., North Potomac, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES

DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

Utility

LEGAL REPRESENTATIVE: HU

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 23
EXEMPLARY CLAIM: 1
LINE COUNT: 16023

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 10 OF 221 USPATFULL on STN

Use of bioactive glass compositions to stimulate osteoblast production Compositions comprising bioactive glass compositions or extracts thereof which include ions in an appropriate concentration and ratio that they enhance osteoblast production, and methods of preparation and use thereof, are disclosed. The compositions can be included in implantable devices that are capable of inducing tissue formation in autogeneic, allogeneic and xenogeneic implants, for example as coatings and/or matrix materials. Examples of such devices include prosthetic implants, sutures, stents, screws, plates, tubes, and the like. Aqueous extracts of the bioactive glass compositions, which extracts are capable of stimulating osteoblast production, are also disclosed. The compositions can be used, for example, to induce local tissue formation from a progenitor cell in a mammal, for accelerating allograft repair in a mammal, for promoting in vivo integration of an implantable prosthetic device to enhance the bond strength between the prosthesis and the existing target tissue at the joining site, and for treating tissue degenerative conditions.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:13078 USPATFULL

TITLE:

Use of bioactive glass compositions to stimulate

osteoblast production

INVENTOR (S):

Hench, Larry L, London, UNITED KINGDOM Polak, Julia M, London, UNITED KINGDOM Buttery, Lee D.k., London, UNITED KINGDOM

Xynos, Ioannis D, Nafplion, GREECE

Maroothynaden, Jason, London, UNITED KINGDOM

NUMBER KIND DATE

US 2004009598 PATENT INFORMATION:

A1 20040115 A1 20030707 US 2003-332731 20030707 (10) APPLICATION INFO.:

WO 2001-US21801 20010711

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

BURNS DOANE SWECKER & MATHIS L L P, POST OFFICE BOX LEGAL REPRESENTATIVE:

1404, ALEXANDRIA, VA, 22313-1404

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 LINE COUNT: 1301

AΒ

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 11 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ТT

The present invention relates to novel polynucleotides associated with the plasma membrane, the polypeptides encoded by these polynucleotides herein collectively referred to as "plasma membrane associated antigens," and antibodies that immunospecifically bind these polypeptides, and the use of such plasma membrane associated polynucleotides, antigens, and antibodies for detecting, treating, preventing and/or prognosing disorders related to these novel polypeptides. More specifically, isolated nucleic acid molecules are provided encoding novel plasma membrane associated polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing these plasma membrane associated polynucleotides, polypeptides, and/or antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the novel polypeptides of the invention. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The invention further relates to methods and/or compositions for inhibiting or promoting the production and/or function of the polypeptides of the invention.

2004:12971 USPATFULL ACCESSION NUMBER:

Nucleic acids, proteins, and antibodies TITLE:

Birse, Charles E., North Potomac, MD, UNITED STATES INVENTOR(S): Rosen, Craig A., Laytonsville, MD, UNITED STATES

> NUMBER KIND DATE _____

US 2004009491 A1 20040115 US 2002-264237 A1 20021004 (10) APPLICATION INFO.:

Continuation-in-part of Ser. No. WO 2001-US16450, filed RELATED APPLN. INFO.:

on 18 May 2001, PENDING

NUMBER DATE ______

US 2000-205515P 20000519 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 2.4 EXEMPLARY CLAIM: 1 LINE COUNT: 18144

PATENT INFORMATION:

L11 ANSWER 12 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies

The present invention relates to novel musculoskeletal system related AΒ polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "musculoskeletal system antigens," and the use of such musculoskeletal system antigens for detecting disorders of

the musculoskeletal system, particularly the presence of cancer and cancer metastases. More specifically, isolated musculoskeletal system associated nucleic acid molecules are provided encoding novel musculoskeletal system associated polypeptides. Novel musculoskeletal system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human musculoskeletal system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the musculoskeletal system, including cancer of musculoskeletal tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:12968 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES, 20850 (U.S. corporation)

KIND NUMBER ______

PATENT INFORMATION: APPLICATION INFO.:

PATENT ASSIGNEE(S):

US 2004009488 A1 20040115 US 2002-242515 A1 20020913 A1 20020913 (10)

DATE

Continuation of Ser. No. US 2001-764877, filed on 17 RELATED APPLN. INFO.:

> NUMBER ______

Jan 2001, PENDING

PRIORITY INFORMATION:

US 2000-179065P US 2000-180628P 20000131 (60) 20000204 (60) US 2000-214886P 20000628 (60) US 2000-217487P 20000711 (60) US 2000-225758P US 2000-220963P US 2000-217496P 20000814 (60) 20000726 (60) 20000711 (60) US 2000-225447P 20000814 (60) US 2000-218290P 20000714 (60) US 2000-225757P 20000814 (60) US 2000-226868P 20000822 (60) US 2000-216647P 20000707 (60) US 2000-2156477 US 2000-225267P US 2000-216880P US 2000-225270P US 2000-251869P US 2000-235834P 20000814 (60) 20000707 (60) 20000814 (60) 20001208 (60) 20000927 (60) US 2000-234274P 20000921 (60) US 2000-234223P 20000921 (60) US 2000-228924P 20000830 (60) US 2000-224518P 20000814 (60) US 2000-236369P 20000929 (60) US 2000-224519P 20000814 (60) US 2000-220964P 20000726 (60) US 2000-241809P 20001020 (60) US 2000-249299P 20001117 (60) US 2000-236327P 20000929 (60) US 2000-241785P 20001020 (60) 20001101 (60) US 2000-244617P

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Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

24

LINE COUNT: 32038

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 13 OF 221 USPATFULL on STN

TI Methods for the treatment of carcinoma

The invention concerns compositions and methods for the diagnosis and treatment of neoplastic cell growth and proliferation in mammals, including humans. The invention is based upon the identification of genes that are amplified in the genome of tumor cells, such as renal cell carcinoma. Such gene amplification is expected to be associated with the overexpression of the gene product as compared to normal cells of the same tissue type and contribute to tumorigenesis. Accordingly, the proteins encoded by the amplified genes are believed to be useful targets for the diagnosis and/or treatment (including prevention) of certain cancers, such as renal cell carcinoma, and may act as predictors

of the prognosis of tumor treatment. The present invention is directed to novel methods of diagnosing and treating tumor, such as renal cell carcinoma or Wilms tumor.

ACCESSION NUMBER:

2004:12653 USPATFULL

TITLE:

INVENTOR(S):

Methods for the treatment of carcinoma

Gerritsen, Mary E., San Mateo, CA, UNITED STATES

Peale, Franklin V., JR., San Carlos, CA, UNITED STATES

Wu, Thomas D., San Francisco, CA, UNITED STATES

PATENT ASSIGNEE(S):

GENENTECH, INC. (U.S. corporation)

KIND DATE NUMBER _______

PATENT INFORMATION:

APPLICATION INFO.:

US 2004009171 A1 20040115 US 2003-372683 A1 20030221 (10)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 2002-271690, filed

on 16 Oct 2002, PENDING

NUMBER

DATE ______

PRIORITY INFORMATION:

US 2001-344534P 20011018 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

GENENTECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA,

94080

NUMBER OF CLAIMS:

57

AB

1 6662

EXEMPLARY CLAIM: LINE COUNT:

L11 ANSWER 14 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies TI

The present invention relates to novel ovarian related polynucleotides, the polypeptides encoded by these polynucleotides herein collectively referred to as "ovarian antigens," and antibodies that immunospecifically bind these polypeptides, and the use of such ovarian polynucleotides, antigens, and antibodies for detecting, treating, preventing and/or prognosing disorders of the reproductive system, particularly disorders of the ovaries and/or breast, including, but not limited to, the presence of ovarian and/or breast cancer and ovarian and/or breast cancer metastases. More specifically, isolated ovarian nucleic acid molecules are provided encoding novel ovarian polypeptides. Novel ovarian polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human ovarian polynucleotides, polypeptides, and/or antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the ovaries and/or breast, including ovarian and/or breast cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The invention further relates to methods and/or compositions for inhibiting or promoting the production and/or function of the polypeptides of the invention.

ACCESSION NUMBER:

2004:7345 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Birse, Charles E., North Potomac, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES

NUMBER								K	Ι	N	D		DATE														
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PATENT INFORMATION: US 2004005579 A1 20040108 APPLICATION INFO.: US 2002-264049 A1 20021004 (10)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. WO 2001-US18569, filed

on 7 Jun 2001, PENDING

NUMBER DATE -----

PRIORITY INFORMATION:

US 2000-209467P 20000607 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

24 1

18130

EXEMPLARY CLAIM: LINE COUNT:

L11 ANSWER 15 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI

The present invention relates to novel proteins. More specifically, isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:7343 USPATFULL

TITLE:

AΒ

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

Human Genome Sciences, Inc., Rockville, MD, UNITED STATES (U.S. corporation)

KIND NUMBER _____

PATENT INFORMATION: APPLICATION INFO.:

PATENT ASSIGNEE(S):

(10)

20000927 (60)

RELATED APPLN. INFO.:

US 2004005577 A1 20040108 US 2002-242747 A1 20020913 Continuation of Ser. No. US 2001-764881, filed on 17

Jan 2001, PENDING

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Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24
EXEMPLARY CLAIM: 1
LINE COUNT: 27694

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 16 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies

TI AB

The present invention relates to novel cardiovascular system related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "cardiovascular system antigens," and the use of such cardiovascular system antigens for detecting disorders of the cardiovascular system, particularly the presence of cancer of cardiovascular system tissues and cancer metastases. More specifically, isolated cardiovascular system associated nucleic acid molecules are provided encoding novel cardiovascular system associated polypeptides. Novel cardiovascular system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human cardiovascular system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the cardiovascular system, including cancer of cardiovascular system tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

INVENTOR(S):

2004:7341 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES, 20850 (U.S. corporation)

STATES, 20850

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

PATENT ASSIGNEE(S):

US 2004005575 A1 20040108 US 2002-227577 A1 20020826 (10)

Continuation of Ser. No. US 2002-91504, filed on 7 Mar 2002, PENDING Continuation of Ser. No. US 2001-764869,

20000927 (60)

20000921 (60)

20000921 (60)

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filed on 17 Jan 2001, ABANDONED

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Utility
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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

24 1

EXEMPLARY CLAIM: LINE COUNT: 28742

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 17 OF 221 USPATFULL on STN L11

Functional MRI agents for cancer imaging TI

The invention relates to novel magnetic resonance imaging contrast AΒ agents for imaging cancer.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:4285 USPATFULL

TITLE:

INVENTOR(S):

Functional MRI agents for cancer imaging Meade, Thomas J., Altadena, CA, United States Fraser, Scott, La Canada, CA, United States

Jacobs, Russell, Arcadia, CA, United States

PATENT ASSIGNEE(S):

Research Corporation Technologies, Inc., Tucson, AZ,

United States (U.S. corporation)

NUMBER KIND _____ ___

PATENT INFORMATION:

US 6673333 B1

20040106

US 2000-715859 APPLICATION INFO.:

20001117 (9)

NUMBER DATE

PRIORITY INFORMATION:

US 2000-201816P 20000504 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

GRANTED

PRIMARY EXAMINER: LEGAL REPRESENTATIVE: Hartley, Michael G. Dorsey & Whitney LLP, Silva, Robin M., Kossiak, Renee

Μ.

NUMBER OF CLAIMS:

10

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

7 Drawing Figure(s); 5 Drawing Page(s)

LINE COUNT:

2422

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 18 OF 221 USPATFULL on STN

50 human secreted proteins TТ

The present invention relates to novel human secreted proteins and AB · isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2004:2568 USPATFULL

TITLE: INVENTOR(S):

50 human secreted proteins Moore, Paul A., Germantown, MD, UNITED STATES Ruben, Steven M., Olney, MD, UNITED STATES

LaFleur, David W., Washington, DC, UNITED STATES Shi, Yanggu, Gaithersburg, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES Olsen, Henrik S., Gaithersburg, MD, UNITED STATES Ebner, Reinhard, Gaithersburg, MD, UNITED STATES Brewer, Laurie A., St. Paul, MN, UNITED STATES

PATENT ASSIGNEE(S):

Human Genome Sciences, Inc., Rockville, MD (U.S.

corporation)

KIND DATE NUMBER ______

PATENT INFORMATION:

APPLICATION INFO.:

RELATED APPLN. INFO.:

US 2004002591 A1 20040101 US 2002-47021 A1 20020117 (10) Continuation-in-part of Ser. No. US 2000-722329, filed

on 28 Nov 2000, PENDING Continuation of Ser. No. US

1999-262109, filed on 4 Mar 1999, ABANDONED

Continuation-in-part of Ser. No. WO 1998-US18360, filed

on 3 Sep 1998, PENDING

NUMBER DATE PRIORITY INFORMATION: US 2001-262066P 20010118 (60)

19970905 (60) US 1997-57626P

US 1997-57663P 19970905 (60)

US 1997-57669P 19970905 (60)

US 1997-58666P 19970912 (60)

US 1997-58667P 19970912 (60)

19970912 (60) US 1997-58973P

19970912 (60) US 1997-58974P

US 1998-90112P 19980622 (60)

Utility DOCUMENT TYPE:

APPLICATION FILE SEGMENT:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 23 EXEMPLARY CLAIM:

2 Drawing Page(s) NUMBER OF DRAWINGS:

33379 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 19 OF 221 USPATFULL on STN

Novel human gene relating to respiratory diseases, obesity, and ΤI

inflammatory bowel disease

This invention relates to genes identified from human chromosome AB 20p13-p12, which are associated with various diseases, including asthma. The invention also relates to the nucleotide sequences of these genes, isolated nucleic acids comprising these nucleotide sequences, and isolated polypeptides or peptides encoded thereby. The invention further relates to vectors and host cells comprising the disclosed nucleotide sequences, or fragments thereof, as well as antibodies that bind to the encoded polypeptides or peptides. Also related are ligands that modulate the activity of the disclosed genes or gene products. In addition, the invention relates to methods and compositions employing the disclosed nucleic acids, polypeptides or peptides, antibodies, and/or ligands for use in diagnostics and therapeutics for asthma and other diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

2004:2447 USPATFULL ACCESSION NUMBER:

Novel human gene relating to respiratory diseases, TITLE:

obesity, and inflammatory bowel disease

Keith, Tim, Bedford, MA, UNITED STATES INVENTOR (S):

Little, Randall D., Newtonville, MA, UNITED STATES

Eerdewegh, Paul Van, Weston, MA, UNITED STATES

Dupuis, Josee, Newton, MA, UNITED STATES

Del Mastro, Richard G., Norfolk, MA, UNITED STATES

Simon, Jason, Westfield, NJ, UNITED STATES Allen, Kristin, Hopkinton, MA, UNITED STATES Pandit, Sunil, Gaithersburg, MD, UNITED STATES

NUMBER	KIND	DATE

PATENT INFORMATION: US 2004002470 A1 20040101

US 2002-277216 A1 20021017 (10) APPLICATION INFO.:

Continuation-in-part of Ser. No. US 2002-126022, filed RELATED APPLN. INFO.:

on 19 Apr 2002, PENDING Continuation-in-part of Ser. No. US 2001-834597, filed on 13 Apr 2001, PENDING Continuation-in-part of Ser. No. US 2000-548797, filed

on 13 Apr 2000, PENDING

Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT:

MORGAN & FINNEGAN, L.L.P., 345 PARK AVENUE, NEW YORK, LEGAL REPRESENTATIVE:

NY, 10154

NUMBER OF CLAIMS: 45 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 162 Drawing Page(s)

LINE COUNT: 15810

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 20 OF 221 USPATFULL on STN

TI Detection and modulation of Slit and roundabount (Robo) mediated

angiogenesis and uses thereof

AB This invention is generally in the field of methods for diagnosis, treatment and prevention of various disorders involving the Slit2

mediated angiogenesis.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:335332 USPATFULL

TITLE:

Detection and modulation of Slit and roundabount (Robo)

mediated angiogenesis and uses thereof

INVENTOR(S):

Geng, Jian-Guo, Portage, MI, UNITED STATES

NUMBER DATE

PRIORITY INFORMATION:

US 2002-362485P 20020308 (60)

DOCUMENT TYPE:

Utility APPLICATION

FILE SEGMENT: LEGAL REPRESENTATIVE:

Peng Chen, Morrison & Foerster LLP, Suite 500, 3811

Valley Centre Drive, San Diego, CA, 92130-2332

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: 1

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

4 Drawing Page(s)

LINE COUNT.

AB

1337

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 21 OF 221 USPATFULL on STN

TI Nucleic acids, proteins, and antibodies

The present invention relates to novel excretory system related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "excretory system antigens," and the use of such excretory system antigens for detecting disorders of the excretory system, particularly the presence of cancer of excretory system tissues and cancer metastases. More specifically, isolated excretory system associated nucleic acid molecules are provided encoding novel excretory system associated polypeptides. Novel excretory system polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human excretory system associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the excretory system, including cancer of excretory system tissues, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:334955 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES

PATENT ASSIGNEE(S):

Human Genome Sciences, Inc., Rockville, MD, 20850 (U.S. corporation)

DATE

	NUMBER	KIND	DATE		
PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:	US 2003235831 US 2002-242355 Continuation of Jan 2001, PENDIN	A1 Ser. No	20031225 20020913 . US 2001-	(10) 764897,	filed on 17

NUMBER

PRIORITY INFORMATION:

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DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24
EXEMPLARY CLAIM: 1
LINE COUNT: 22457

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 22 OF 221 USPATFULL on STN

TI Nucleic acids, proteins, and antibodies

The present invention relates to novel proteins. More specifically, isolated nucleic acid molecules are provided encoding novel polypeptides. Novel polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human polynucleotides and/or polypeptides, and antibodies. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to these novel polypeptides. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting or enhancing the production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER: 2003:334953 USPATFULL

TITLE: Nucleic acids, proteins, and antibodies INVENTOR(S): Ruben, Steven M., Olney, MD, UNITED STATES

Barash, Steven C., Rockville, MD, UNITED STATES
Rosen, Craig A., Laytonsville, MD, UNITED STATES
Birse, Charles E., North Potomac, MD, UNITED STATES
Human Conome Sciences, Inc., Rockville, MD, UNITED

PATENT ASSIGNEE(S): Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES (U.S. corporation)

PATENT INFORMATION: US 2003235829 A1

APPLICATION INFO.: US 2002-227646 A1 20020826 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-860670, filed on 21 May 2001, PENDING Continuation-in-part of Ser. No. WO

2001-US1346, filed on 17 Jan 2001, PENDING

NUMBER DATE

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DOCUMENT TYPE:

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

1

Utility

EXEMPLARY CLAIM: LINE COUNT: 20415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 23 OF 221 USPATFULL on STN

Compositions and methods for systemic inhibition of cartilage TI degradation

Methods and compositions for inhibiting articular cartilage degradation. AΒ The compositions preferably include multiple chondroprotective agents, including at least one agent that promotes cartilage anabolic activity and at least one agent that inhibits cartilage catabolism. The compositions may also include one or more pain and inflammation

inhibitory agents. The compositions may be administered systemically, such as to treat patients at risk of cartilage degradation at multiple joints, and suitably may be formulated in a carrier or delivery vehicle that is targeted to the joints. Alternatively the compositions may be injected or infused directly into the joint.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

2003:334713 USPATFULL ACCESSION NUMBER:

Compositions and methods for systemic inhibition of TITLE:

cartilage degradation

Demopulos, Gregory A., Mercer Island, WA, UNITED STATES INVENTOR(S):

Palmer, Pamela Pierce, San Francisco, CA, UNITED STATES

Herz, Jeffrey M., Mill Creek, WA, UNITED STATES

Omeros Corporation (U.S. corporation) PATENT ASSIGNEE(S):

KIND DATE NUMBER -----US 2003235589 A1 20031225 US 2003-356649 A1 20030131 PATENT INFORMATION: APPLICATION INFO.: (10) Continuation-in-part of Ser. No. US 2002-31546, filed RELATED APPLN. INFO.:

on 18 Jan 2002, PENDING A 371 of International Ser. No.

WO 2000-US19864, filed on 21 Jul 2000, PENDING

Continuation-in-part of Ser. No. US 2001-839633, filed on 20 Apr 2001, PENDING Continuation-in-part of Ser. No. WO 1999-US26330, filed on 5 Nov 1999, PENDING Continuation-in-part of Ser. No. WO 1999-US24625, filed

on 20 Oct 1999, PENDING

DATE NUMBER -----PRIORITY INFORMATION: US 2002-353552P 20020201 (60) 19990721 (60) US 1999-144904P US 1998-107256P 19981105 (60) US 1998-105026P 19981020 (60)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

OMEROS MEDICAL SYSTEMS, INC., 1420 FIFTH AVENUE, SUITE LEGAL REPRESENTATIVE:

2675, SEATTLE, WA, 98101

NUMBER OF CLAIMS: 155 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 9 Drawing Page(s)

LINE COUNT: 6575

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 24 OF 221 USPATFULL on STN

Nucleic acids, proteins, and antibodies ΤI AB The present invention relates to novel endocrine related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "endocrine antigens," and the use of such endocrine antigens for detecting disorders of the endocrine system, particularly the presence of cancers of the endocrine system and endocrine cancer metastases. More specifically, isolated endocrine associated nucleic acid molecules are provided encoding novel endocrine associated polypeptides. Novel endocrine polypeptides and antibodies that bind to these polypeptides are provided. Also provided are vectors, host cells, and recombinant and synthetic methods for producing human endocrine associated polynucleotides and/or polypeptides. The invention further relates to diagnostic and therapeutic methods useful for diagnosing, treating, preventing and/or prognosing disorders related to the endocrine system, including cancers of the endocrine system, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of polynucleotides and polypeptides of the invention. The present invention further relates to methods and/or compositions for inhibiting the

production and function of the polypeptides of the present invention.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:330759 USPATFULL

TITLE:

Nucleic acids, proteins, and antibodies

INVENTOR (S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Ruben, Steven M., Olney, MD, UNITED STATES Barash, Steven C., Rockville, MD, UNITED STATES Human Genome Sciences, Inc., Rockville, MD (U.S.

PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE ______

PATENT INFORMATION:

US 2003232975 A1 20031218 US 2002-74024 A1 20020214

APPLICATION INFO.: RELATED APPLN. INFO.:

(10) Continuation of Ser. No. US 2001-764895, filed on 17

Jan 2001, ABANDONED

		NUMBER	DATE	
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US 2001-259678P
Utility
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DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

24

LINE COUNT: 21828

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 25 OF 221 USPATFULL on STN

TТ Proteases

The invention provides human proteases (PRTS) and polynucleotides which AB identify and encode PRTS. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with aberrant expression of PRTS.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ACCESSION NUMBER:

2003:330138 USPATFULL

TITLE:

Proteases

INVENTOR(S):

Delegeane, Angelo M., Milpitas, CA, UNITED STATES Gandhi, Ameena R., San Francisco, CA, UNITED STATES Hafalia, April J. A., Santa Clara, CA, UNITED STATES Lu, Dyung Aina M., San Jose, CA, UNITED STATES Arvizu, Chandra S., San Jose, CA, UNITED STATES Tribouley, Catherine M., San Francisco, CA, UNITED

STATES Das, Debopriya, Mountain View, CA, UNITED STATES Kallick, Deborah A., Portola Valley, CA, UNITED STATES Nguyen, Danniel B., San Jose, CA, UNITED STATES Lee, Ernestine A., Castro Valley, CA, UNITED STATES Khan, Farrah A., Glen View, IL, UNITED STATES Yue, Henry, Sunnyvale, CA, UNITED STATES Au-Young, Janice, Brisbane, CA, UNITED STATES Griffin, Jennifer A., Fremont, CA, UNITED STATES Policky, Jennifer L., San Jose, CA, UNITED STATES

Ramkumar, Jayalaxmi, Fremont, CA, UNITE Yang, Junming, San Jose, CA, UNITED STATE Thangavelu, Kavitha, Mountain View, CA, UN STATES Ding, Li, Creve Coeur, MO, UNITED STATES Kearney, Liam, San Francisco, CA, UNITED ST Baughn, Mariah R., San Leandro, CA, UNITED STATES Borowsky, Mark L., Redwood City, CA, UNITED STATES Sanjanwala, Madhusudan, Los Altos, CA, UNITED STATES Yao, Monique G., Carmel, IN, UNITED STATES Burford, Neil, Durham, CT, UNITED STATES Chawla, Narinder K., Union City, CA, UNITED STATES Lal, Preeti G., Santa Clara, CA, UNITED STATES Lee, Sally, San Jose, CA, UNITED STATES Todd, Stephen, San Francisco, CA, UNITED STATES Lo, Terence P., Foster City, CA, UNITED STATES Tang, Y. Tom, San Jose, CA, UNITED STATES Elliott, Vicki S., San Jose, CA, UNITED STATES Azimzai, Yalda, Oakland, CA, UNITED STATES Lu, Yan, Palo Alto, CA, UNITED STATES

PATENT ASSIGNEE(S):

Incyte Genomics, Inc., Palo Alto, CA (U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION:

US 2003232349 A1 20031218 US 2002-274639 A1 20021018

APPLICATION INFO.: RELATED APPLN. INFO.:

PRIORITY INFORMATION:

US 2002-274639 Al 20021018 (10) Continuation of Ser. No. WO 2001-US22397, filed on 17

Jul 2001, PENDING

NUMBER DATE
-----US 2000-220063P 20000721 (60)
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Utility APPLICATION

LEGAL REPRESENTATIVE:

INCYTE CORPORATION (formerly known as Incyte, Genomics,

Inc.), 3160 PORTER DRIVE, PALO ALTO, CA, 94304

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CAS INDEXING IS AVAILABLE FOR THIS PATENT.